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<td>September 30, 2019</td>
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<td>September 16, 2019</td>
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<td>March 31, 2020</td>
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<td>February 18, 2020</td>
<td>March 16, 2020</td>
<td>March 31, 2020</td>
<td>April 30, 2020</td>
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Documents will be accepted for filing on all regular workdays from 8:00 a.m. until 5:00 p.m. We encourage early filings to facilitate the timely publication of the Missouri Register. Orders of Rulemaking appearing in the Missouri Register will be published in the Code of State Regulations and become effective as listed in the chart above. Advance notice of large volume filings will facilitate their timely publication. We reserve the right to change the schedule due to special circumstances. Please check the latest publication to verify that no changes have been made in this schedule. To review the entire year’s schedule, please check out the website at sos.mo.gov/adrules/pubsched.
HOW TO CITE RULES AND RSMO

RULES
The rules are codified in the *Code of State Regulations* in this system–

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<thead>
<tr>
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<th>Chapter</th>
<th>Rule</th>
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<td>3</td>
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<td>10-</td>
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<tr>
<td>Department</td>
<td>Agency</td>
<td>General area regulated</td>
<td>Specific area regulated</td>
</tr>
<tr>
<td><em>Code of State Regulations</em></td>
<td>Division</td>
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</tr>
</tbody>
</table>

and should be cited in this manner: 3 CSR 10-4.115.

Each department of state government is assigned a title. Each agency or division in the department is assigned a division number. The agency then groups its rules into general subject matter areas called chapters and specific areas called rules. Within a rule, the first breakdown is called a section and is designated as (1). Subsection is (A) with further breakdown into paragraphs 1., subparagraphs A., parts (I), subparts (a), items I. and subitems a.

The rule is properly cited by using the full citation, for example, 3 CSR 10-4.115 NOT Rule 10-4.115.

Citations of RSMo are to the *Missouri Revised Statutes* as of the date indicated.

*Code and Register on the Internet*

The *Code of State Regulations* and *Missouri Register* are available on the Internet.

The *Code* address is sos.mo.gov/adrules/csr/csr

The *Register* address is sos.mo.gov/adrules/moreg/moreg

These websites contain rulemakings and regulations as they appear in the *Code* and *Registers*. 
Rules appearing under this heading are filed under the authority granted by section 536.025, RSMo. An emergency rule may be adopted by an agency if the agency finds that an immediate danger to the public health, safety, or welfare, or a compelling governmental interest requires emergency action; follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances; follows procedures which comply with the protections extended by the *Missouri* and the *United States Constitutions*; limits the scope of such rule to the circumstances creating an emergency and requiring emergency procedure, and at the time of or prior to the adoption of such rule files with the secretary of state the text of the rule together with the specific facts, reasons, and findings which support its conclusion that there is an immediate danger to the public health, safety, or welfare which can be met only through the adoption of such rule and its reasons for concluding that the procedure employed is fair to all interested persons and parties under the circumstances.

All emergency rules must state the period during which they are in effect, and in no case can they be in effect more than one hundred eighty (180) calendar days or thirty (30) legislative days, whichever period is longer. Emergency rules are not renewable, although an agency may at any time adopt an identical rule under the normal rulemaking procedures.

Title 12—DEPARTMENT OF REVENUE  
Division 10—Director of Revenue  
Chapter 41—General Tax Provisions  

**EMERGENCY AMENDMENT**

12 CSR 10-41.010 Annual Adjusted Rate of Interest. The department proposes to amend the purpose, emergency statement, section (1), and authority.

**PURPOSE:** This emergency amendment establishes the annual adjusted rate of interest to be implemented and applied on taxes remaining unpaid during calendar year 2020.

**EMERGENCY STATEMENT:** The director of revenue is mandated to establish not later than October 22 annual adjusted rate of interest based upon the adjusted prime rate charged by banks during September of that year as set by the Board of Governors of the Federal Reserve rounded to the nearest full percent. This emergency amendment is necessary to ensure public awareness and to preserve a compelling governmental interest requiring an early effective date in that the amendment informs the public of the established rate of interest to be paid on unpaid amounts of taxes for the 2020 calendar year. A proposed amendment, that covers the same material, is published in this issue of the Missouri Register. The director has limited the scope of the emergency amendment to the circumstances creating the emergency. The director has followed procedures calculated to assure fairness to all interested persons and parties and has complied with protections extended by the *Missouri* and *United States Constitutions*. Emergency amendment filed October 21, 2019, becomes effective January 1, 2020, expires June 28, 2020.

(1) Pursuant to section 32.065, RSMo, the director of revenue upon official notice of the average predominant prime rate quoted by commercial banks to large businesses, as determined and reported by the Board of Governors of the Federal Reserve System in the Federal Reserve Statistical Release H.15(519) for the month of September of each year has set by administrative order the annual adjusted rate of interest to be paid on unpaid amounts of taxes during the succeeding calendar year as follows:

<table>
<thead>
<tr>
<th>Calendar Year</th>
<th>Rate of Interest on Unpaid Amounts of Taxes</th>
</tr>
</thead>
<tbody>
<tr>
<td>1995</td>
<td>12%</td>
</tr>
<tr>
<td>1996</td>
<td>9%</td>
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<tr>
<td>1997</td>
<td>8%</td>
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<td>9%</td>
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<td>10%</td>
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<td>2002</td>
<td>6%</td>
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<td>2003</td>
<td>5%</td>
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<td>2004</td>
<td>4%</td>
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<td>2018</td>
<td>4%</td>
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<tr>
<td>2019</td>
<td>5%</td>
</tr>
<tr>
<td>2020</td>
<td>5%</td>
</tr>
</tbody>
</table>


**PUBLIC COST:** This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

**PRIVATE COST:** This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.
I. RULE NUMBER

<table>
<thead>
<tr>
<th>Rule Number and Name:</th>
<th>12 CSR 10-41.010 Annual Adjusted Rate of Interest</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type of Rulemaking:</td>
<td>Emergency Amendment</td>
</tr>
</tbody>
</table>

II. SUMMARY OF FISCAL IMPACT

<table>
<thead>
<tr>
<th>Affected Agency or Political Subdivision</th>
<th>Estimated Cost of Compliance in the Aggregate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Counties</td>
<td>This proposed amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate. The 2020 interest rate imposed on delinquent taxes will be the same as the rate imposed in 2019.</td>
</tr>
<tr>
<td>Cities</td>
<td></td>
</tr>
<tr>
<td>Special Taxing Districts</td>
<td></td>
</tr>
</tbody>
</table>

III. WORKSHEET

The proposed amendment establishes the rate of interest for 2020 at five percent (5%), remaining the same as the rate in 2019.

The future amount of past due taxes is unknown. With the 2020 interest rate imposed upon delinquent taxes remaining the same as that imposed in 2019, public entities realize no additional fiscal impact. This proposed amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

Interest on Delinquent Taxes Paid to Department of Revenue

<table>
<thead>
<tr>
<th></th>
<th>Current Rule 5.00%</th>
<th>Proposed Amendment 5.00%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Past due tax amount</td>
<td>$100.00</td>
<td>$100.00</td>
</tr>
<tr>
<td>Interest Amount (%)</td>
<td>x 5.00</td>
<td>x 5.00</td>
</tr>
<tr>
<td>Total Amount Due</td>
<td>$105.00</td>
<td>$105.00</td>
</tr>
</tbody>
</table>
IV. ASSUMPTIONS
Pursuant to Section 32.065, RSMo, the Director of Revenue is mandated to establish an annual adjusted rate of interest based upon the adjusted prime rate charged by banks during September of that year, as set by the Board of Governors of the Federal Reserve, rounded to the nearest full percentage. The actual bank prime loan rate noted by the Federal Reserve in 2019 was 5.25 percent.
FISCAL NOTE
PRIVATE COST

I. RULE NUMBER

Rule Number and Name: 12 CSR 10-41.010 Annual Adjusted Rate of Interest
Type of Rulemaking: Emergency Amendment

II. SUMMARY OF FISCAL IMPACT

<table>
<thead>
<tr>
<th>Estimate of the number of entities by class which would likely be affected by adoption of the proposed rule:</th>
<th>Classification by types of the business entities which would likely be affected:</th>
<th>Estimate in the aggregate as to the cost of compliance with the rule by the affected entities:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any taxpayer with delinquent tax.</td>
<td>Any taxpayer with delinquent tax.</td>
<td>This proposed amendment will not cost private entities more than five hundred dollars ($500) in the aggregate. The 2020 interest rate imposed on delinquent taxes remains the same as that imposed in 2019. The actual number of affected taxpayers is unknown.</td>
</tr>
</tbody>
</table>

III. WORKSHEET

The proposed amendment establishes the rate of interest for 2020 at five percent (5%), the same as the rate in 2019.

This proposed amendment will not cost private entities more than five hundred dollars ($500) in the aggregate. Because the 2020 interest rate imposed on delinquent taxes remains at the same rate as that imposed in 2019, the interest rate remains the same on each $100 of delinquent taxes to private entities. The actual number of affected taxpayers is unknown.

Interest on Delinquent Taxes Paid to Department of Revenue

<table>
<thead>
<tr>
<th>Current Rule</th>
<th>Proposed Amendment</th>
</tr>
</thead>
<tbody>
<tr>
<td>5.00%</td>
<td>5.00%</td>
</tr>
<tr>
<td>Past due tax amount</td>
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</tr>
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<td>Interest Amount (%)</td>
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<tr>
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</tbody>
</table>

IV. ASSUMPTIONS

Pursuant to Section 32.065, RSMo, the Director of Revenue is mandated to establish an annual adjusted rate of interest based upon the adjusted prime rate charged by banks during September of that year, as set by the Board of
Governors of the Federal Reserve, rounded to the nearest full percentage. The actual bank prime loan rate noted by the Federal Reserve in 2019 was 5.25 percent.
Emergency Amendment

13 CSR 70-10.030 Prospective Reimbursement Plan for Nonstate-Operated Facilities for ICF/IID Services

The division is amending sections (2), (3), (4), (7), (8), and (10), deleting section (6), and renumbering the remaining sections.

PURPOSE: The MO HealthNet Divisions seeks through this amendment to rebase the per diem rates paid to nonstate-operated intermediate care facilities for individuals with intellectual disabilities (ICF/IID) for services, to clarify the process for determining reimbursement rates, to remove and/or replace obsolete processes and language, and to combine and remove duplicative language.

EMERGENCY STATEMENT: The Department of Social Services, MO HealthNet Division, by rule and regulation, must define the reasonable costs, manner, extent, quantity, quality, charges, and fees of medical assistance provided to MO HealthNet participants. Effective for dates of service beginning January 1, 2019, the MO HealthNet Division will rebases the per diem rates of nonstate-operated ICF/IID facilities using a more current year cost report base. The appropriation by the General Assembly for State Fiscal Year (SFY) 2019 and SFY 2020 included additional funds to increase nonstate-operated ICF/IID reimbursement rates which will be used to fund the rebased per diem rates. The increased reimbursement resulting from the rebased rates is necessary to ensure that payments for ICF/IID per diem rates are in line with the funds appropriated for that purpose. There are a total of seven (7) nonstate-operated ICF/IID providers currently enrolled in Missouri Medicaid, all of which will receive a rebased per diem rate. This emergency amendment will ensure payment for ICF/IID services to approximately seventy-nine (79) ICF/IID Missourians in accordance with the appropriation authority. For the rebased per diem rates to be implemented, the MO HealthNet Division was required to submit a Medicaid State Plan Amendment (SPA) to the Centers for Medicare and Medicaid Services (CMS). CMS approved the SPA on June 24, 2019. The proposed state regulation will be effective on or around April 30, 2020. This emergency amendment must be implemented on a timely basis to ensure that quality ICF/IID services continue to be provided to Medicaid patients in ICF/IID facilities in accordance with the appropriation authority. As a result, the MO HealthNet Division finds an immediate danger to public health, safety and/or welfare and a compelling governmental interest, which requires emergency action. The Missouri Medical Assistance program has a compelling government interest in providing continued cash flow for ICF/IID services and to adequately compensate these providers for costs expended on the state Medicaid population that they serve. The scope of this emergency amendment is limited to the circumstances creating the emergency and complies with the protections extended by the Missouri and United States Constitutions. The MO HealthNet Division believes that this emergency amendment is fair to all interested persons and parties under the circumstances. A proposed amendment covering this same material will be published in the Missouri Register. This emergency amendment was filed October 25, 2019, becomes effective November 8, 2019, and expires May 5, 2020.

(2) General Principles.

(A) The MO HealthNet program shall reimburse qualified providers of ICF/IID services based solely on the individual MO HealthNet participant’s days of care (within benefit limitations) multiplied by the facility’s Title XIX per diem rate less any payments made by participants.

(B) Effective November 1, 1986, the Title XIX per diem rate for all ICF/IID facilities participating on or after October 31, 1986, shall be the lower of—

1. The average private pay charge;
2. The Medicare per diem rate, if applicable;
3. The rate paid to a facility on October 31, 1986, as adjusted by updating its base year to its 1985 fiscal year. Facilities which do not have a full twelve- (12-) month 1985 fiscal year shall not have their base years updated to their 1985 fiscal years. Changes in ownership, management, control, operation, leasehold interests by whatever form for any facility previously certified for participation in the MO HealthNet program at any time that results in increased capital costs for the successor owner, management, or leaseholder shall not be recognized for purposes of reimbursement; and

4. However, any provider who does not have a rate on October 31, 1986, and whose facility meets the definition in subsection (3)(J) of this rule, will be exempt from paragraph (2)(B)3., and the rate shall be determined in accordance with applicable provisions of this rule.

1. The Medicare per diem rate, if applicable; or
2. The reimbursement rate as determined in accordance with this regulation.

(C) This plan has an effective date of November 1, 1986, at which time prospective per diem rates shall be calculated for the remainder of the state’s FY-87 and future fiscal years. Per diem rates established by updating facilities’ base years to FY-85 may be subject to retroactive and prospective adjustment based on audit of the facilities’ new base year period.

(D) The Title XIX per diem rates as determined by this plan shall apply only to services furnished on or after November 1, 1986.

(E) All illustrations and examples provided throughout this rule are for illustration purposes only and are not meant to be actual calculations.

(3) Definitions.

(A) “Allowable cost areas” means those cost areas which are allowable for allocation to the MO HealthNet program based upon the principles established in this rule. The allowable cost areas, not specifically addressed in this rule, will be based upon criteria of the Medicare Provider Reimbursement Manual (HM-15) and section 17(6) of this rule.

(B) “Average private pay charge” means the average private pay charge for non-MO HealthNet patients determined by dividing total non-MO HealthNet days of care into total revenue collected for the same service that is included in the MO HealthNet per diem rate, excluding negotiated payment methodologies with the Veterans Administration and the Missouri Department of Mental Health.

(C) Committee. The advisory committee defined in subsection (6)(A) of this rule.

(D) “Cost report” means the cost report detailing the cost of rendering covered services for the fiscal reporting period. Providers must file the cost report on timely forms provided by and in accordance with the procedures of the Department of Social Services.

(E) “Departmental. The department, unless otherwise specified, refers to” means the Missouri Department of Social Services, unless otherwise specified.

(F) “Director. The director, unless otherwise specified, refer to” means the director of the Missouri Department of Social Services, unless otherwise specified.

(G) “Effective date” means November 1, 1986.

2. The effective date for rate adjustments granted in accordance with section (6) of this rule shall be for dates of service beginning the first day of the month following the director’s, or his/her designee’s, final determination on the rate.
(H) (G) “ICF/IID. Nonstate-operated” means nonstate-operated facilities certified to provide intermediate care for individuals with intellectual disabilities under the Title XIX program.

(I) (H) “Medicare rate. This is Rate” means the allowable cost of care permitted by Medicare standards and principles of reimbursement.

(J) (I) “New [construction. Newly] Construction means newly built facilities or parts, for which an approved Certificate of Need (CON) or applicable waivers were obtained and which were newly completed and operational on or after November 1, 1986.


(L) (K) “Providers. A provider” means, under the Prospective Reimbursement Plan [is], a nonstate-operated ICF/IID facility with a valid participation agreement, in effect on or after October 31, 1986, with the Missouri Department of Social Services for the purpose of providing long-term care (LTC) services to Title XIX-eligible participants. Facilities certified to provide intermediate care services to individuals with intellectual disabilities under the Title XIX program may be offered a MO HealthNet participation agreement on or after January 1, 1990, only if (1) the facility has no more than fifteen (15) beds for individuals with intellectual disabilities, and (2) there is no other licensed residential living facility for individuals with intellectual disabilities within a radius of one-half (1/2) mile of the facility seeking participation in the MO HealthNet program.

(M) (L) “Reasonable and [adequate reimbursement. Reimbursement] Adequate Reimbursement” means reimbursement levels which meet the needs of an efficiently and economically operated facility and which in no case exceed normal market costs.

(N) (M) “Related parties. Parties are related when —” means—

1. An individual or group, regardless of the business structure of either, where, through their activities, one (1) individual’s or group’s transactions are for the benefit of the other and the benefits exceed those which are usual and customary in the dealings;

2. One (1) or more persons [has] have an ownership or controlling interest in a party, and the person(s) or one (1) or more relatives of the person(s) has an ownership or controlling interest in the other party. For the purposes of this paragraph, ownership or controlling interest does not include a bank, savings bank, trust company, building and loan association, savings and loan association, credit union, industrial loan and thrift company, investment banking firm, or insurance company unless the entity, directly or through a subsidiary, operates a facility; or

3. As used in section (3), the following terms mean:

A. “Indirect [ownership/interest] Ownership” or “Indirect Interest” means an ownership interest in an entity that has an ownership interest in another entity. This term includes an ownership interest in any entity that has an indirect ownership interest in an entity;

B. “Ownership [interest] Interest” means the possession of equity in the capital, in the stock, or in the profits of an entity;

C. “Ownership for controlling interest is when” Interest” or “Controlling Interest” means a person or corporation(s)—

(I) Has an ownership interest at the total totalizing five percent (5%) or more in an entity;

(II) Has an indirect ownership interest equal to five percent (5%) or more in an entity. The amount of indirect ownership interest is determined by multiplying the percentages of ownership in each entity;

(III) Has a combination of direct and indirect ownership interest equal to five percent (5%) or more in an entity;

(IV) Owns an interest of five percent (5%) or more in any mortgage, deed of trust, note, or other obligation secured by an entity, if that interest equals at least five percent (5%) of the value of the property or assets of the entity. The percentage of ownership resulting from the obligations is determined by multiplying the percentage of interest owned in the obligation by the percentage of the entity’s assets used to secure the obligation;

(V) Is an officer or director of an entity; or

(VI) Is a partner in an entity that is organized as a partnership;

D. “Relative” means persons related by blood or marriage to the fourth degree of consanguinity; and

E. “Entity” means any person, corporation, partnership, or association.

(O) (N) “Rural. Those” means those counties [which] that are not defined as urban.

(P) (O) “Urban. The urban counties are” means counties that are standard metropolitan statistical areas including Andrew, Boone, Buchanan, Cass, Christian, Clay, Franklin, Greene, Jackson, Jasper, Jefferson, Newton, Platte, Ray, St. Charles, St. Louis, and St. Louis City.

(4) [Prospective Reimbursement] ICF/IID Rate Computation. Except in accordance with other provisions of this rule, the provisions of this section shall apply to all providers of ICF/IID services certified to participate in Missouri’s MO HealthNet program. Rate determination shall be based on reasonable and adequate reimbursement levels for allowable cost items described in this rule which are related to ordinary and necessary care for the level-of-care provided for an efficiently and economically operated facility. All providers shall submit documentation of expenses for allowable cost areas. The department shall have authority to require those uniform accounting and reporting procedures and forms as it deems necessary. A reasonable and adequate reimbursement in each allowable cost area will be determined.

A. [Except in accordance with other provisions of this rule, the provisions of this section shall apply to all providers of ICF/IID services certified to participate in Missouri’s MO HealthNet program.] Prospective Reimbursement Rate Determination through December 31, 2018.

1. ICF/IID facilities

A. Except in accordance with other provisions of this rule, the MO HealthNet program shall reimburse providers of LTC services based on the individual MO HealthNet-participating days of care multiplied by the Title XIX prospective per diem rate less any payments collected from participants. The Title XIX prospective per diem reimbursement rate for the remainder of state Fiscal Year 1987 shall be the facility’s per diem reimbursement payment rate in effect on October 31, 1986, as adjusted by updating the facility’s allowable base year to its 1985 fiscal year. Each facility’s per diem costs as reported on its Fiscal Year 1985 Title XIX cost report will be determined in accordance with the principles set forth in this rule. If a facility has not filed a 1985 fiscal year cost report, the MO HealthNet Division will use the most current cost report on file with the department [will be used] to set its/a facility’s per diem rate. Facilities with less than a full twelve-(12)-month 1985 fiscal year will not have their base year rates updated.

B. For state FY-88 and dates of service beginning July 1, 1987, the negotiated trend factor shall be equal to two percent (2%) to be applied in the following manner: Two percent (2%) of the average per diem rate paid to both state- and nonstate-operated ICF/IID facilities on June 1, 1988, shall be added to each facility’s rate.

C. For state FY-89 and dates of service beginning January 1, 1989, the negotiated trend factor shall be equal to one percent (1%) to be applied in the following manner: One percent (1%) of the average per diem rate paid to both state- and nonstate-operated ICF/IID facilities on June 1, 1988, shall be added to each facility’s rate.

D. For state FY-91 and dates of service beginning July 1, 1990, the negotiated trend factor shall be equal to one percent (1%) to be applied in the following manner: One percent (1%) of the average per diem rate paid to both state- and nonstate-operated ICF/IID facilities on June 1, 1990, shall be added to each facility’s rate.
5. Prospective payment adjustment (PPA). A FY92 PPA will be provided prior to the end of the state fiscal year for nonstate-operated ICF/IID facilities with a current provider agreement on file with the MO HealthNet Division as of October 1, 1991.

A. For providers that qualify, the PPA shall be the lesser of—

   (I) The provider’s facility peer group factor (FPGF) times the projected patient days (PPD) covered by the adjustment year times the prospective payment adjustment factor (PPAF) times the nonstate-operated intermediate care facility for individuals with intellectual disabilities ceiling (ICFIIIDC) on October 1, 1991 (FPGF × PPD × PPAF × ICFIIIDC). For example: A provider having nine hundred twenty (920) paid days for the period May 1991 to July 1991 out of a total paid days for this same period of twenty-eight thousand five hundred sixty-one (28,561) represents an FPGF of three and twenty-two hundredths percent (3.22%). So using the FPGF of 3.22% × 114,244 × 24.5% × $156.01 = $140,607; or

   (II) The provider FPGF times one hundred forty-five percent (145%) of the amount credited to the intermediate care revenue collection center (ICRCC) of the State Title XIX Fund (STF) for the period October 1, 1991 through December 31, 1991.

B. FPGF—is determined by using each ICF/IID facility’s paid days for the service dates in May 1991 through July 1991 as of September 20, 1991, divided by the sum of the paid days for the same service dates for all providers qualifying as of the determination date of October 16, 1991.

C. ICFIIIDC—is one hundred fifty-six dollars and one cent ($156.01) on October 1, 1991.

D. PPAF—is equal to twenty-four and one half percent (24.5%) for fiscal year 1992 which includes an adjustment for economic trends.

E. PPD—is the projection of one hundred fourteen thousand two hundred forty-four (114,244) patient days made on October 1, 1991, for the adjustment year.

6. FY-92 trend factor and Workers’ Compensation. All facilities with either an interim rate or a prospective per diem rate in effect on September 1, 1992, shall be granted an increase to their per diem rate effective September 1, 1992, of eight dollars and eighty-six cents ($8.86) per patient day related to the continuation of the FY-92 trend factor and the Workers’ Compensation adjustment. This adjustment is equal to seven and one-half percent (7.5%) of the March 1992 weighted average per diem rate of one hundred eighty dollars and fourteen cents ($180.14) for all nonstate-operated ICF/IID facilities.

7. FY-93 negotiated trend factor. All facilities with either an interim rate or prospective per diem rate in effect on September 1, 1992, shall be granted an increase to their per diem rate effective September 1, 1992, of one dollar and sixty-six cents ($1.66) per patient day for the negotiated trend factor. This adjustment is equal to one and four-tenths percent (1.4%) of the March 1992 weighted average per diem rate of one hundred eighty dollars and fourteen cents ($180.14) for all nonstate-operated ICF/IID facilities.

E. FPA. FY-96 negotiated trend factor. All nonstate-operated ICF/IID facilities shall be granted an increase to their per diem rates effective for dates of service beginning January 1, 1996, of sixty dollars and seven cents ($60.07) per patient day for the negotiated trend factor. This adjustment is equal to three percent (3%) of the weighted average per diem rate paid to nonstate-operated ICF/IID facilities on June 30, 1998, of one hundred forty-eight dollars and ninety-nine cents ($148.99).

G. FY-2000 trend factor. All nonstate-operated ICF/IID facilities shall be granted an increase to their per diem rates effective for dates of service beginning January 1, 1999, of four dollars and sixty-three cents ($4.63) per patient day for the trend factor. This adjustment is equal to three percent (3%) of the weighted average per diem rate paid to nonstate-operated ICF/IID facilities on April 30, 1999, of one hundred fifty-four dollars and forty-three cents ($154.43). This increase shall only be used for increases for the salaries and fringe benefits for direct care staff and their immediate supervisors.

H. FY-2001 trend factor. All nonstate-operated ICF/IID facilities shall be granted an increase to their per diem rates effective for dates of service beginning July 1, 2000, of forty dollars and eighty-one cents ($40.81) per patient day for the trend factor. This adjustment is equal to four hundred eighty-six thousand two hundred forty-four (86,244) patient days made on October 1, 2000, of one hundred sixty dollars and twenty-three cents ($160.23). This increase shall only be used for increases for the salaries and fringe benefits for direct care staff and their immediate supervisors.

I. FY-2002 trend factor. All nonstate-operated ICF/IID facilities shall be granted an increase of seven percent (7%) to their per diem rates effective for dates of service billed for state fiscal year 2002 and thereafter. This adjustment is equal to one percent (1%) of the per diem rate paid to nonstate-operated ICF/IID facilities on June 30, 2002.

J. FY-2003 trend factor. Effective for dates of service beginning July 1, 2003, all nonstate-operated ICF/IID facilities shall be granted an increase to their per diem rates of two percent (2%) for the trend factor. This adjustment is equal to two percent (2%) of the per diem rate paid to nonstate-operated ICF/IID facilities on June 30, 2003.

K. FY-2004 trend factor. Effective for dates of service beginning July 1, 2004, all nonstate-operated ICF/IID facilities shall be granted an increase to their per diem rates of three percent (3%) for the trend factor. This adjustment is equal to three percent (3%) of the per diem rate paid to nonstate-operated ICF/IID facilities on June 30, 2004.

L. FY-2005 catch up increase. Effective for dates of service beginning July 1, 2005, all nonstate-operated ICF/IID facilities shall be granted an increase to their per diem rates of five percent (5%) for the trend factor. This adjustment is equal to five percent (5%) of the per diem rate paid to nonstate-operated ICF/IID facilities on June 30, 2005.

M. FY-2006 trend factor. Effective for dates of service beginning July 1, 2006, all nonstate-operated ICF/IID facilities shall be granted an increase to their per diem rates of seven percent (7%) for the trend factor. This adjustment is equal to seven percent (7%) of the per diem rate paid to nonstate-operated ICF/IID facilities on June 30, 2006.

N. FY-2007 trend factor. Effective for dates of service beginning July 1, 2007, all nonstate-operated ICF/IID facilities shall be granted an increase to their per diem rates of one percent (1%) for the trend factor. This adjustment is equal to one percent (1%) of the per diem rate paid to nonstate-operated ICF/IID facilities on June 30, 2007.

O. FY-2008 trend factor. Effective for dates of service beginning July 1, 2008, all nonstate-operated ICF/IID facilities shall be granted an increase to their per diem rates of one percent (1%) for the trend factor. This adjustment is equal to one percent (1%) of the per diem rate paid to nonstate-operated ICF/IID facilities on June 30, 2008.

P. FY-2009 trend factor. Effective for dates of service beginning July 1, 2009, all nonstate-operated ICF/IID facilities shall be granted an increase to their per diem rates of one percent (1%) for the trend factor. This adjustment is equal to one percent (1%) of the per diem rate paid to nonstate-operated ICF/IID facilities on June 30, 2009.
beginning September 1, 2016, all nonstate-operated ICF/IID facilities shall be granted an increase to their per diem rates of two percent (2%) for the trend factor. This adjustment is equal to two percent (2%) of the per diem rate paid to nonstate-operated ICF/IID facilities on August 31, 2016.

/O.20/ State FY-2018 per diem adjustment. Effective for dates of service beginning September 1, 2017, all nonstate-operated ICF/IID facilities shall be subject to a decrease to their per diem rates of two and eighty-two hundredths percent (2.82%). This adjustment is equal to two and eighty-two hundredths percent (2.82%) of the per diem rate paid to nonstate-operated ICF/IID facilities on August 31, 2017.

(B) Per Diem Rate Calculation Effective for Dates of Service Beginning January 1, 2019. Effective for dates of service beginning January 1, 2019, the MO HealthNet Division shall rebase nonstate-operated ICF/IID facilities’ per diem rates using the facilities’ 2017 fiscal year end cost reports. The rebased rates are contingent upon approval of the state plan amendment by the Centers for Medicare and Medicaid Services.

1. Prospective Rate Calculation.

A. Each nonstate-operated ICF/IID shall have its prospective rate recalculated based on its 2017 fiscal year end cost report using the same principles and methodology as detailed throughout sections (1)-(13) of this regulation.

I. The total costs from the 2017 fiscal year end cost reports shall be trended using the indices from the most recent publication of the Healthcare Cost Review Available to the division using the “CMS Nursing Home without Capital Market Basket” table. The costs shall be trended using the four quarter moving average. The costs shall be trended for the years following the cost report year, up to and including the state fiscal year corresponding to the effective date of the rates. For SFY 2019, the trends are as follows:

   (a) 2018 = 3.025%
   (b) 2019 = 2.65%

II. If a facility’s total calculated per diem set forth in this section is less than the facility’s current rate, the facility shall continue to receive its current rate.

III. The division will use the FY 2017 cost report to determine the ICF/IID prospective rate, set forth as follows:

(a) Total Routine Service Cost. Total routine service cost includes patient care, ancillary, dietary, laundry, housekeeping, plant operations, and administration. Each ICF/IID’s Title XIX Routine Service Cost per diem shall be calculated as follows:

1. The total routine service costs as reported on the cost report shall be adjusted for minimum utilization, if applicable, trended to the current state fiscal year, and divided by the total patient days to determine the per diem. The minimum utilization adjustment will be determined by applying the unused capacity percent to the sum of the laundry, housekeeping, plant operations, and administration expenses. The following is an illustration of how this item (4)(B)1.A. (III) (a) is calculated:

| Licensed/Certified Bed Days (9 beds x 365 days) | 3,285 |
| Total Patient Days | 2,900 |
| Percent Occupied (2,900/3,285) | 88% |
| Bed Days @ Minimum Occupancy of 90% (3,285 x 90%) | 2,957 |
| Unused Capacity (90% of Bed Days Less Total Patient Days) | 57 |
| Unused Capacity Percent for Minimum Utilization | 1.93% |
| Adjustment (Unused Capacity / 90% of Bed Days) | 1.93% |
| Minimum Utilization Days for Return on Owner’s Equity (Greater of 90% of Bed Days or Total Patient Days) | 2,957 |

* Minimum Utilization Adjustment

| Laundry | $5,000 |
| Housekeeping | $8,000 |
| Plant Operations | $46,000 |
| Administration | $165,000 |
| Total Expense | $224,000 |
| Unused Capacity Percent | 1.93% |
| Minimum Utilization Adjustment | $4,323 |

Unused Capacity Percent x Total Expense | $4,323 |

Routine Service Cost, Adjusted for Minimum Utilization | $654,677 |

SFY 2018 Trend | 3.025% |

SFY 2019 Trend | 2.65% |

Trended Routine Service Cost | $692,355 |

Total Patient Days | 2,900 |

Routine Service Cost Per Diem | $238.74 |

(b) Intermediate Care Facility for Individuals with Intellectual Disabilities Federal Reimbursement Allowance (ICF/IID FRA). The SFY 2019 ICF/IID FRA provider assessment as determined in accordance with 9 CSR 10-31.030 is divided by total patient days to determine the ICF/IID FRA per diem.

1. The following is an illustration of how the ICF/IID FRA assessment is calculated:

| SFY 2019 ICF/IID FRA Assessment | $40,000 |
| Total Patient Days | 2,900 |
| ICF/IID FRA Per Diem | $13.79 |

(c) Return on Equity. An owner’s net equity consists of investment capital and working capital as indicated in subsection (6)(S). Each ICF/IID’s Return on Equity per diem is calculated as follows:

I. Investment Capital. Investment capital includes the investment in building, property and equipment (cost of land, mortgage payments toward principal and equipment purchase less the accumulated depreciation).

II. Working Capital. Working capital represents the amount of capital which is required to ensure proper operation of the facility and shall be calculated as 1.1 months of the total expenses less depreciation.

III. The total net equity shall be multiplied by the rate of return as set forth in Section (6)(S) to determine the return on equity. The return on equity is subject to the minimum occupancy percent of 90% in determining the per diem.

IV. The following is an illustration of how this item (4)(B)1.A. (III) (c) is calculated:

<table>
<thead>
<tr>
<th>Investment Capital</th>
<th>Equipment</th>
<th>Building</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cost</td>
<td>$130,000</td>
<td>$300,000</td>
<td>$430,000</td>
</tr>
<tr>
<td>Less: Prior Years Depreciation</td>
<td>($120,000)</td>
<td>($225,000)</td>
<td>($345,000)</td>
</tr>
<tr>
<td>Less: Current Year Depreciation</td>
<td>($2,400)</td>
<td>($8,500)</td>
<td>($10,900)</td>
</tr>
<tr>
<td>Total Investment Capital</td>
<td>$7,600</td>
<td>$66,500</td>
<td>$74,100</td>
</tr>
</tbody>
</table>
Emergency Rules

Working Capital
Total Expenses $659,000
Less: Current Year Depreciation Expense ($10,900) $648,100
Divided by 12 Months $54,008
Times 1.1 Months $59,409
Total Working Capital

Net Equity (Investment Capital + Working Capital) $133,509
Rate of Return 5.125%
Return on Equity $6,842
Working Capital) $133,509
Net Equity (Investment Capital + Working Capital) $133,509
Rate of Return 5.125%
Return on Equity $6,842
Minimum Utilization Days 2,957
Return on Equity Per Diem $2.31

(c) Rebased Per-Diem Rate. The total calculated Per-Diem is the sum of the Routine Service Cost per diem, the ICF/IID FRA per diem and the Return on Equity per diem. To determine the rebased per diem rate, the total calculated per diem is compared to the current per diem rate and the facility will be held harmless if the total calculated per diem is less than the current per diem rate (i.e., if the total calculated per diem is less than the current per diem rate, the facility would receive the current per diem).

<table>
<thead>
<tr>
<th>Routine Service Cost per diem</th>
<th>$238.74</th>
</tr>
</thead>
<tbody>
<tr>
<td>ICF/IID FRA per diem</td>
<td>$13.79</td>
</tr>
<tr>
<td>Return on Equity per diem</td>
<td>$2.31</td>
</tr>
<tr>
<td>Total Calculated Per Diem</td>
<td>$254.84</td>
</tr>
</tbody>
</table>

Current Per Diem Rate $200.00

Rebased Per Diem Rate $254.84

(If the total calculated per diem is less than the current per diem rate, the facility would receive the current per diem rate)

2. Interim Rate Calculation.
A. In the case of a newly certified facility where a valid Title XIX participation agreement has been executed, a request for an interim rate must be submitted in writing to the MO HealthNet Division.

(I) The interim rate shall be determined based on the projected estimated operating costs. The facility’s request must specifically and clearly identify the interim rate and be supported by complete and accurate documentation satisfactory to the single state agency. Documentation submitted must include a budget of the projected estimated operating costs. Other documentation may also be required to be submitted upon the request of the division.

(II) The establishment of the prospective rate for all new construction facility providers shall be based on the second full facility fiscal year cost report (i.e., rate setting cost report) prepared in accordance with the principles of this rule. This cost report shall be based on actual operating costs and shall be prepared and submitted in accordance with the reporting requirements in section (7) of this rule.

(III) Prior to establishment of a prospective rate for newly certified facility providers, the cost reports may be subject to an on-site audit by the Department of Social Services or authorized representative to determine the facility’s actual allowable costs. Allowability of costs will be determined as described in subsection (3)(A) of this rule.

(IV) The cost report, audited or unaudited, will be reviewed by the MO HealthNet Division, and a prospective reimbursement rate shall be determined on the allowable per diem cost as set forth in section (4) of this rule. The prospective reimbursement rate shall be effective on the first day of the facility’s rate setting cost report and payment adjustments shall be made for claims paid at the interim rate.

[2./3. Adjustments to rates. The prospectively determined reimbursement rate may be adjusted only under the following conditions:
A. When information contained in a facility’s cost report is found to be fraudulent, misrepresented, or inaccurate, the facility’s reimbursement rate may be reduced, both retroactively and prospectively, if the fraudulent, misrepresented, or inaccurate information as originally reported resulted in establishment of a higher reimbursement rate than the facility would have received in the absence of this information. No decision by the MO HealthNet agency to impose a rate adjustment in the case of fraudulent, misrepresented, or inaccurate information in any way shall affect the MO HealthNet agency’s ability to impose any sanctions authorized by statute or rule. The fact that fraudulent, misrepresented, or inaccurate information reported did not result in establishment of a higher reimbursement rate than the facility would have received in the absence of the information also does not affect the MO HealthNet agency’s ability to impose any sanctions authorized by statute or rules;
B. In accordance with subsection (6)(B) of this rule, a newly constructed facility’s initial reimbursement rate may be reduced if the facility’s actual allowable per diem cost for its first twelve (12) months of operation is less than its initial rate;
C. When a facility’s MO HealthNet reimbursement rate is higher than either its private pay rate or its Medicare rate, the MO HealthNet rate will be reduced in accordance with subsection (2)(B) of this rule/ Extraordinary circumstances. A participating facility that has a prospective rate may request an adjustment to its prospective rate due to extraordinary circumstances. This request should be submitted in writing to the division within one (1) year of the occurrence of the extraordinary circumstance. The request should clearly and specifically identify the conditions for which the rate adjustment is sought. The dollar amount of the requested rate adjustment should be supported by complete and accurate documentation satisfactory to the division. If the division makes a written request for additional information and the facility does not comply within ninety (90) days of the request for additional information, the division shall consider the request withdrawn. Requests for rate adjustments that have been withdrawn by the facility or are considered withdrawn because of failure to supply requested information may be resubmitted once for the requested rate adjustment. In the case of a rate adjustment request that has been withdrawn and then resubmitted, the effective date shall be the first day of the month in which the resubmitted request was made providing that it was made prior to the tenth day of the month. If the resubmitted request is not filed by the tenth of the month, rate adjustments shall be effective the first day of the following month. Conditions for an extraordinary circumstance are as follows:

[(D.)](I) When the provider can show that it incurred higher costs due to circumstances beyond its control, and the circumstances are not experienced by the nursing home or ICF/IID industry in general, and the [request must] circumstances have a substantial cost effect. These circumstances include, but are not limited to;

[(II)](a) Unavoidable [A]acts of nature, such as/ are natural wildfire, earthquakes, hurricane, tornado, lightning, and flood/ flooding, or other natural disasters for which no one can be held responsible, that are not covered by insurance and that occur in a federally declared disaster area; or

[(II)](b) Vandalism, civil disorder, or both that are not covered by insurance; or

[(III)](c) Replacement of capital depreciable items not built into existing rates that are the result of circumstances not related to normal wear and tear or upgrading of existing system;]
E. When an adjustment to a facility’s rate is made in accordance with the provisions of section (6) of this rule; or
F. When an adjustment is based on an Administrative Hearing Commission or court decision.

D. New, expanded, or terminated services may be subject to rate review.

E. Disallowance of federal financial participation.

F. The following will not be subject to review:
   (I) The negotiated trend factor;
   (II) The use of prospective reimbursement rate; and
   (III) The cost base for the per diem rates except as specified in this rule.

(18) In the case of newly constructed nonstate-operated ICF/IID facilities entering the MO HealthNet program after October 31, 1986, and for which no rate has previously been set, the director or his/her designee may set an initial rate for the facility as in his/her discretion s/he deems appropriate. The initial rate shall be subject to review by the advisory committee under the provisions of section (6) of this rule.

(5) Covered Services and Supplies.

(A) ICF/IID services and supplies covered by the per diem reimbursement rate under this plan, and which the ICF/IID must provide, as required by federal or state law or rule and include, among other services, the regular room, dietary and nursing services, or any other services that are required for standards of participation or certification. Also included are minor medical and surgical supplies and the use of equipment and facilities. These items include, but are not limited to, the following:

1. All general nursing services including, but not limited to, administration of oxygen and related medications, hand-feeding, incontinency care, tray service, and enemas;

2. Items [which] that are furnished routinely and relatively uniformly to all participants, for example, gowns, water pitchers, soap, basins, and bed pans;

3. Items such as alcohol, applicators, cotton balls, band-aids, and tongue depressors;

4. All nonlegend antacids, nonlegend laxatives, nonlegend stool softeners, and nonlegend vitamins. Any nonlegend drug in one of these four (4) categories must be provided to residents as needed and no additional charge may be made to any party for any of these drugs. Facilities may not elect which nonlegend drugs in any of the four (4) categories to supply; [all must be provided] facilities must provide all as needed within the existing per diem rate;

5. Items which are utilized by individual participants but which are reusable and expected to be available, such as ice bags, bed rails, canes, crutches, walkers, wheelchairs, traction equipment, and other durable, nondepreciable medical equipment;

6. Additional items as specified in the appendix to this plan when required by the patient;

7. Special dietary supplements used for tube feeding or oral feeding, such as elemental high nitrogen diet, including dietary supplements written as a prescription item by a physician;

8. All laundry services except personal laundry, which is a non-covered service;

9. All general personal care services [which are furnished] that the facility furnishes routinely and relatively uniformly to all participants for their personal cleanliness and appearance shall be covered services, for example, necessary clipping and cleaning of fingernails and toenails, basic hair care, shampoos, and shaves to the extent necessary for reasonable personal hygiene. The provider shall not bill the patient or his/her responsible party for this type of personal service;

10. All consultative services as required by state or federal law or regulation or for proper operation by the provider. Contracts for the purchase of these services must accompany the provider cost report. Failure to do so will result in the penalties specified in section [(9)/(8) of this rule;]

11. Semiprivate room and board and private room and board when necessary to isolate a participant due to a medical or social condition, such as contagious infection, irrational loud speech, and the like. Unless a private room is necessary due to a medical or social condition, a private room is a noncovered service, and a MO HealthNet participant or responsible party may therefore pay the difference between a facility’s semiprivate charge and its charge for a private room. MO HealthNet participants may not be placed in private rooms and charged any additional amount above the facility’s MO HealthNet per diem unless the participant or responsible party in writing specifically requests a private room prior to placement in a private room and acknowledges that an additional amount not payable by MO HealthNet will be charged for a private room;

12. Twelve (12) days per any period of six (6) consecutive months during which a participant is on a temporary leave of absence from the facility. [Temporary leave of absence days must be specifically provided for] The provider shall specifically provide for temporary leave of absence days in the participant’s plan of care. Periods of time during which a participant is away from the facility because s/he is visiting a friend or relative are considered temporary leaves of absence; and

13. Days when participants are away from the facility overnight on facility-sponsored group trips under the continuing supervision and care of facility personnel.

(6) Rate Determination. All nonstate-operated ICF/IID providers of LTC services under the MO HealthNet program who desire to have their rates changed or established must apply to the MO HealthNet Division. The department may request the participation of the Department of Mental Health in the analysis for rate determination. The procedure and conditions for rate reconsideration are as follows:

(A) Advisory Committee. The director, Department of Social Services, shall appoint an advisory committee to review and make recommendations pursuant to provider requests for rate determination. The director may accept, reject, or modify the advisory committee’s recommendations.

1. Membership. The advisory committee shall be composed of four (4) members representative of the nursing home industry in Missouri, three (3) members from the Department of Social Services, and two (2) members who may include, but are not limited to, a consumer representative, an accountant or economist, or a representative of the legal profession. Members shall be appointed for terms of twelve (12) months. The director shall select a chairman from the membership who shall serve at the director’s discretion.

2. Procedures.
   A. The committee may hold meetings when five (5) or more members are present and may make recommendations to the department in instances where a simple majority of those present and voting concur.
   B. The committee shall meet no less than one (1) time each quarter, and members shall be reimbursed for expenses.
   C. The MO HealthNet Division will summarize each case and, if requested by the advisory committee, make recommendations. The advisory committee may request additional documentation as well as require the facility to submit to a comprehensive operational review to determine if there exists an efficient and economical delivery of patient services. The review will be made at the discretion of the committee and may be performed by it or its designee. The findings from a review may be used to determine the per diem rate for the facility. Failure to submit requested documentation shall be grounds for denial of the request.
D. The committee, at its discretion, may issue its recommendation based on written documentation or may request further justification from the provider sending the request.

E. The advisory committee shall have ninety (90) days from the receipt of each complete request, provided the request is on behalf of a facility which has executed a valid Title XIX participation agreement, or the receipt of any additional documentation to submit its recommendations in writing to the director. If the committee is unable to make a recommendation within the specified time limit, the director or his/her designee, if the committee establishes good cause, may grant a reasonable extension.

F. Final determination on rate adjustment. The director’s, or his/her designee’s, final decision on each request shall be issued in writing to the provider within fifteen (15) working days from receipt of the committee’s recommendation.

G. The director’s, or his/her designee’s, final determination on the advisory committee’s recommendation shall become effective on the first day of the month in which the request was made, providing that it was made prior to the tenth of the month. If the request is not filed by the tenth of the month, adjustments shall be effective the first day of the following month;

(B) In the case of new construction where a valid Title XIX participation agreement has been executed, a request for a rate must be submitted in writing to the MO HealthNet Division and must specifically and clearly identify the issue and the total amount involved. The total dollar amount must be supported by complete, accurate, and documented records satisfactory to the single state agency. Until an initial per diem rate is established, the MO HealthNet Division shall grant a tentative per diem rate for that period. In no case may a facility receive a per diem reimbursement rate greater than the class ceiling in effect on March 1, 1990, adjusted by the negotiated trend factor.

1. In the case of newly built facility or part of the facility which is less than two (2) years of age and enters the Title XIX Program on or after November 1, 1986, a reimbursement rate shall be assigned based on the projected estimated operating costs. Advice of the advisory committee will be obtained for all initial rate determination requests for new construction. Owners of new construction which have an approved CON are certified for participation and which have a valid Title XIX participation agreement shall submit a budget in accordance with the principles of section (7) of this rule and other documentation as the committee may request.

2. The establishment of the permanent rate for all new construction facility providers shall be based on the second full facility fiscal year cost report prepared in accordance with the principles of section (7) of this rule. This cost report shall be submitted within ninety (90) days of the close of the provider’s second full facility fiscal year. This cost report shall be based on actual operating costs. No request for an extension of this ninety- (90-) day filing requirement will be considered. Any new construction facility. Any construction facility provider which fails to timely submit the cost report may be subject to sanction under this rule and 13 CSR 70-3.030.

3. Prior to establishment of a permanent rate for new construction facility providers, the cost reports may be subject to an on-site audit by the Department of Social Services to determine the facility’s actual allowable costs. Allowability of costs will be determined as described in subsection (3)(A) of this rule.

4. The cost report, audited or unaudited, will be reviewed by the MO HealthNet Division, and each facility’s actual allowable per diem cost will be determined. The cost report shall not be submitted to the advisory committee for review. If a facility’s actual allowable per diem cost is less than its initial per diem reimbursement rate, the facility’s rate will be reduced to its actual allowable per diem cost. This reduction will be effective on the first day of the second full facility fiscal year.

5. If a facility’s actual allowable per diem cost is higher than its initial per diem reimbursement rate, the facility’s rate will not be adjusted; a facility shall not receive a rate increase based on review or audit of the cost report and actual operating costs;

(C) In the case of existing facilities not previously certified to participate in the Title XIX program, a request for a per diem reimbursement rate must be submitted in writing to the MO HealthNet Division and must specifically and clearly identify the issue and the total amount involved. The total dollar amount must be supported by complete, accurate, and documented records satisfactory to the single state agency. Until the time as a per diem rate is established, the MO HealthNet Division shall grant a tentative per diem rate for that period. In no case may a facility receive a per diem reimbursement rate greater than the class ceiling in effect on March 1, 1990, adjusted by the negotiated trend factor.

1. In the case of a facility described in subsection (6)(C) of this rule and entering the Title XIX program on or after March 1, 1990, a reimbursement rate shall be assigned based on the projected estimated operating costs. Advice of the advisory committee will be obtained for all initial rate determination requests for first full facility’s fiscal year.

2. The establishment of the permanent rate for all existing facility providers shall be based on the second full facility fiscal year cost report prepared in accordance with the principles of section (7) of this rule. This cost report shall be submitted within ninety (90) days of the close of their second full facility fiscal year. This cost report shall be based on actual operating costs. No request for an extension of this ninety- (90-) day filing requirement will be considered. Any new construction facility provider which fails to timely submit the cost report may be subject to sanction under this rule and 13 CSR 70-3.030.

3. Prior to establishment of a permanent rate for existing facility providers, the cost reports may be subject to an on-site audit by the Department of Social Services to determine the facility’s actual allowable costs. Allowability of costs will be determined as described in subsection (3)(A) of this rule.

4. The cost report, audited or unaudited, will be reviewed by the MO HealthNet Division, and each facility’s actual allowable per diem cost will be determined. The cost report shall not be submitted to the advisory committee for review. If a facility’s actual allowable per diem cost is less than its initial per diem reimbursement rate, the facility’s rate will be reduced to its actual allowable per diem cost. This reduction will be effective on the second day of the first full facility fiscal year.

5. If a facility’s actual allowable per diem cost is higher than its initial per diem reimbursement rate, the facility’s rate will not be adjusted; a facility shall not receive a rate increase based on review or audit of the cost report and actual operating costs;

(D) Rate Reconsideration.

1. The committee may review the following conditions for rate reconsideration:

   A. Those costs directly related to a change in a facility’s case mix; and

   B. Requests for rate reconsideration which the director, in his/her discretion, may refer to the committee due to
The request for an adjustment must be submitted in writing to the MO HealthNet Division and must specifically and clearly identify the issue and the total dollar amount involved. The total dollar amount must be supported by complete, accurate, and documented records satisfactory to the single state agency. The facility must demonstrate that the adjustment is necessary, proper, and consistent with efficient and economical delivery of covered patient care services.

3. However, for state fiscal years after Fiscal Year 1987, in no case may a facility receive a per diem reimbursement rate higher than the class ceiling for that facility in effect on June 30 of the preceding fiscal year adjusted by the negotiated trend factor.

4. The following will not be subject to review:
   A. The negotiated trend factor;
   B. The use of prospective reimbursement rate; and
   C. The cost base for the June 30 per diem rate except as specified in this rule;

(E) Rate Adjustments. The department may alter a facility’s per diem rate based on—
   1. Court decisions;
   2. Administrative Hearing Commission decisions;
   3. Determination through desk audits, field audits, and other means, which establishes misrepresentations in or the inclusion of unallowable costs in the cost report used to establish the per diem rate. In these cases, the adjustment shall be applied retroactively; or
   4. Adjustments determined by the department without the advice of the rate advisory committee.

A. Prospective payment adjustment (PPA). A FY-92 PPA will be provided prior to the end of the state fiscal year for nonstate-operated ICF/IID facilities with a current provider agreement on file with the MO HealthNet Division as of October 1, 1991.

   (I) For providers which qualify, the PPA shall be the lesser of—
   (a) The provider’s facility peer group factor (FPGF) times the projected patient days (PPD) covered by the adjustment year times the prospective payment adjustment factor (PPAF) times the nonstate-operated intermediate care facility for individuals with intellectual disabilities ceiling (ICF/IIDC) on October 1, 1991 (FPGF × PPD × PPAF × ICF/IIDC).
   For example: A provider having nine hundred twenty (920) paid days for the period May 1991 to July 1991 out of a total paid days for this same period of twenty-eight thousand five hundred sixty-one (28,561) represents an FPGF of three and twenty-two hundredths percent (3.22%) times the projected patient days (28,561) times the provider’s facility peer group factor (FPGF) times the nonstate-operated intermediate care facility for individuals with intellectual disabilities ceiling (ICF/IIDC) on October 1, 1991.

   (b) The provider FPGF times one hundred forty-five percent (145%) of the amount credited to the intermediate care revenue collection center (ICRCC) of the State Title XIX Fund (STF) for the period October 1, 1991 through December 31, 1991.

   (II) PPGF—is determined by using each ICF/IID facility’s paid days for the service dates in May 1991 through July 1991 as of September 20, 1991, divided by the sum of the paid days for the same service dates for all provider’s qualifying as of the determination date of October 16, 1991.

   (III) ICF/IIDC—is one hundred fifty-six dollars and one cent ($156.01) on October 1, 1991.

   (IV) PPAF—is equal to twenty-four and five-tenths percent (24.5%) for fiscal year 1992 which includes an adjustment for economic trends.

(V) PPD—is the projection of one hundred fourteen thousand two hundred forty-four (114,244) patient days made on October 1, 1991, for the adjustment year;

4. Adjustments determined by the department in this section shall be applied retroactively; or

5. FY-92 trend factor and Workers’ Compensation. All facilities with either an interim rate or a prospective per diem rate in effect on September 1, 1992, shall be granted an increase to their per diem rate effective September 1, 1992, of eight dollars and eighty-six cents ($8.86) per patient day related to the continuation of the FY-92 trend factor and the Workers’ Compensation adjustment. This adjustment is equal to seven and one-half percent (7.5%) of the March 1992 weighted average per diem rate of one hundred eighteen dollars and fourteen cents ($118.14) for all nonstate-operated ICF/IID facilities; or

6. FY-93 negotiated trend factor. All facilities with either an interim rate or prospective per diem rate in effect on September 1, 1992, shall be granted an increase to their per diem rate effective September 1, 1992, of one dollar and sixty-six cents ($1.66) per patient day for the negotiated trend factor. This adjustment is equal to one and four-tenths percent (1.4%) of the March 1992 weighted average per diem rate of one hundred eighteen dollars and fourteen cents ($118.14) for all nonstate-operated ICF/IID facilities; and

   (F) Rate determination shall be based on a determination of reasonable and adequate reimbursement levels for allowable cost items described in this rule which are related to ordinary and necessary care for the level-of-care provided for an efficiently and economically operated facility. All providers shall submit documentation of expenses for allowable cost areas. The department shall have authority to require those uniform accounting and reporting procedures and forms as it deems necessary. A reasonable and adequate reimbursement in each allowable cost area will be determined by the advisory committee with the consent of the director.

[(7)](6) Allowable Cost Areas.

(A) Compensation of Owners.

1. Allowance of compensation of services of owners shall be an allowable cost area, provided the [services are actually performed] owner actually performs the services and the services are necessary [services].

2. “Compensation” [shall mean] means the total benefit to the owner, within the limitations set forth in this rule, [by the owner] of the services s/he renders to the facility [including]. Compensation includes direct payments to the owner for managerial, administrative, professional, and other services;[;] amounts paid by the provider for the personal benefit of the owner;[;] the cost of assets and services which the owner receives from the provider;[;] and additional amounts determined to be the reasonable value of the services rendered by sole proprietors or partners and not paid by any method previously described.

(B) Covered services and supplies as defined in section (5) of this rule.

(C) Depreciation.

1. An appropriate allowance for depreciation on buildings, furnishings, and equipment [which] are part of the operation and sound conduct of the provider’s business is an allowable cost item.

2. Depreciation is part of the operation and sound conduct of the provider’s business.
FINDER’S FEES ARE NOT AN ALLOWABLE COST ITEM.

2. The depreciation must be identifiable and recorded in the provider’s accounting records, based on the historical cost of the asset and prorated over the estimated useful life of the asset using the straight-line method of depreciation from the date initially put into service.

3. The basis of assets at the time placed in service shall be the lower of—
   A. The book value of the provider;
   B. Fair market value at the time of acquisition;
   C. The recognized Internal Revenue Service (IRS) tax basis; and
   D. In the case of the change in ownership, the cost basis of acquired assets of the owner of record on or after July 18, 1984, as of the effective date of the change of ownership; or in the case of a facility which entered the program after July 18, 1984, the owner at the time of the initial entry into the MO HealthNet program.

4. The MO HealthNet Division will allow the basis of donated assets will be allowed to the extent of recognition of income resulting from the donation of the asset. Should a dispute arise between a provider and the Department of Social Services as to the fair market value at the time of acquisition of a depreciable asset and an appraisal by a third party is required, the appraisal cost will be shared proportionately by the MO HealthNet program and the facility in ratio to MO HealthNet participant reimbursable patient days to total patient days.

5. Allowable methods of depreciation shall be limited to the straight-line method. The depreciation method used for an asset under the MO HealthNet program need not correspond to the method used by a provider for non-MO HealthNet purposes; however, useful life shall be in accordance with the American Hospital Association’s Guidelines. Component part depreciation is optional and allowable under this plan.

6. “Historical cost” [list] means the cost incurred by the provider in acquiring the asset and preparing it for use, except as provided in this rule. Usually, historical cost includes costs that would be capitalized under generally accepted accounting principles. For example, in addition to the purchase price, historical cost would include architectural fees and related legal fees. Where a provider has elected, for federal income tax purposes, to expense certain items such as interest and taxes during construction, the historical cost basis for MO HealthNet depreciation purposes may include the amount of these expensed items. However, where a provider did not capitalize these costs and has written off the costs in the year they were incurred, the provider cannot retroactively capitalize any part of these costs under the program. For Title XIX purposes and this rule, any asset costing less than five hundred dollars ($500) or having a useful life of one (1) year or less, may be expensed and not capitalized at the option of the provider, or in the case of a facility which entered the program after July 18, 1984, the owner at the time of the initial entry into the MO HealthNet program.

7. When an asset is acquired by trading in an existing asset, the cost basis of the new asset shall be the sum of the depreciated cost basis of the traded asset plus the cash paid.

8. For the purpose of determining allowable depreciation, the cost basis of the asset shall be as prescribed in paragraph (7)(6)(C).

9. Capital expenditures for building construction or for renovation costs which are in excess of one hundred fifty thousand dollars ($150,000) and which cause an increase in a provider’s bed capacity shall not be allowed in the program or depreciation base if these capital expenditures fail to comply with any other federal or state law or regulation, such as Certificate of Need (CON).

10. Amortization of leasehold rights and related interest and finance costs shall not be allowable costs under this plan.

(D) Interest and Finance Costs.

1. Necessary and proper interest on both current and capital indebtedness shall be an allowable cost item excluding finder’s fees.

2. Interest is the cost incurred for the use of borrowed funds. Interest on current indebtedness is the cost incurred for funds borrowed for a relatively short term. This is usually for those purposes as working capital for normal operating expenses. Interest on capital indebtedness is the cost incurred for funds borrowed for capital purposes, such as the acquisition of facilities and capital improvements, and this indebtedness must be amortized over the life of the loan.

3. Interest may be included in finance charges imposed by some lending institutions or it may be a prepaid cost or discount in transactions with those lenders who collect the full interest charges when funds are borrowed.

4. To be an allowable cost item, interest (including finance charges, prepaid costs, and discounts) must be supported by evidence of an agreement that funds were borrowed and that payment of interest and repayment of the funds are required, identifiable in the provider’s accounting records, relating to the reporting period in which the costs are claimed, and necessary and proper for the operation, maintenance, or acquisition of the provider’s facilities.

5. Necessary means that the interest be incurred for a loan made to satisfy a financial need of the provider and for a purpose related to participant care. Loans which result in excess funds or investments are not considered necessary.

6. Proper means that the interest be incurred at a rate not in excess of what a prudent borrower would have had to pay in the money market existing at the time the loan was made, and provided further the department shall not reimburse for interest and finance charges any amount in excess of the prime rate current at the time the loan was obtained.

7. Interest on loans to providers by proprietors, partners, and any stockholders shall not be an allowable cost item because the loans shall be treated as invested capital and included in the computation of an allowable return on owner’s net equity. If a facility operated by a religious order borrows from the order, interest paid to the order shall be an allowable cost.

8. If loans for capital indebtedness exceed the asset cost basis as defined in subsection (7)(6)(C) of this rule, the interest associated with the portion of the loan(s) which exceed the asset cost basis as defined in subsection (7)(6)(C) of this rule shall not be allowable.

9. Income from a provider’s qualified retirement fund shall be excluded in consideration of the per diem rate.

10. A provider shall amortize finance charges, prepaid interest, and discount over the period of the loan ratably or by means of the constant rate of interest method on the unpaid balance.

11. Usual and customary costs, excluding finder’s fees, incurred to obtain loans shall be treated as interest expense and shall be allowable costs over the loan period ratably or by means of the constant interest applied method.

12. Usual and customary costs shall be limited to the lender’s title and recording fees, appraisal fees, legal fees, escrow fees, and closing costs.

13. Interest expense resultant from capital expenditures for building construction or for renovation costs which are in excess of one hundred fifty thousand dollars ($150,000) and which cause an increase in a bed capacity by the provider shall not be an allowable cost item if the capital expenditure fails to comply with other federal or state law or rules such as CON.

(E) Rental and Leases.

1. Rental and leases of land, buildings, furnishings, and equipment are allowable cost areas provided that if the rented items are necessary and not in essence a purchase of those assets. Finder’s fees are not an allowable cost item.

2. Necessary rental and lease items are those which are pertinent to the economical operation of the provider.

3. In the case of related parties, rental and lease amounts cannot exceed the lesser of those which are actually paid or the costs to the related party.

4. Determination of reasonable and adequate reimbursement for rental and amounts, except in the case of related parties which are subject to other provisions of this rule, may require affidavits of
competent, impartial experts who are familiar with the current rentals and leases.

5. The test of necessary costs shall take into account the agreement between the owner and the tenant regarding the payment of related property costs.

6. Leases subject to CON approval must have that approval before a rate is determined.

7. If rent or lease costs increase solely as a result of change in ownership, the resulting increase which exceeds the allowable capital cost of the owner of record as of July 18, 1984, or in the case of a facility which entered the program after July 18, 1984, the owner at the time of the initial entry into the MO HealthNet program, shall be a nonallowable cost.

(F) Taxes. Taxes levied on or incurred by providers shall be allowable cost areas with the exceptions of the following items:

1. Federal, state, or local income and excess profit taxes including any interest and penalties paid;

2. Taxes in connection with financing, refinancing, or refunding operations, such as taxes on the issuance of bond, property transfer, issuance of transfer of stocks;

3. Taxes for which exceptions are available to the provider;

4. Special assessments on land [which] represent capital improvements. These costs shall be capitalized and depreciated over the period during which the assessment is scheduled to be paid;

5. Taxes on property which are not a part of the operation of the provider;

6. Taxes which are levied against a resident and collected and remitted by the provider; and

7. Self-employment Federal Insurance Contributions Act (FICA) taxes applicable to individual proprietors, partners, or members of a joint venture to the extent the taxes exceed the amount which would have been paid by the provider on the allowable compensation of the persons had the provider organization been an incorporated rather than unincorporated entity.

(G) Issuance of Revenue Bond and Tax Levies by District and County Facilities. Those nursing home districts and county facilities whose funding is through the issuance of revenue bonds, that interest which is paid per the revenue bond will be an allowable cost item. Depreciation on the plant and equipment of these facilities also shall be an allowable cost item. Any tax levies which are collected by nursing home districts or county homes that are supported in whole or in part by these levies will not be recognized as a revenue offset except to the extent that the funds are used for the actual operation of the facility.

(H) Value of Services of Employees.

1. Except as provided for in this rule, the value of services performed by employees in the facility shall be included as an allowable cost area to the extent actually compensated, either to the employee or to the supplying organization.

2. Services rendered by volunteers, such as those affiliated with the American Red Cross, hospital guilds, auxiliaries, private individuals, and similar organizations, shall not be included as an allowable cost area, as the services have traditionally been rendered on a purely volunteer basis without expectation of any form of reimbursement by the organization through which the service is rendered or by the person rendering the service.

3. Services by priests, ministers, rabbis, and similar type professionals shall be an allowable cost area; provided, that the services are not of a religious nature. An example of an allowable cost area under this section would be a necessary administrative function performed by a clergyman. The state will not recognize building costs on space set aside primarily for professionals providing any religious function. [Costs] The MO HealthNet Division considers costs for wardrobe and similar items likewise [are considered] nonallowable.

(I) Fringe Benefits.

1. Life insurance.

A. Types of insurance [which are not considered] that the MO HealthNet Division does not consider an allowable cost area; premiums related to insurance on the lives of officers and key employees are not allowable cost areas under the following circumstances:

(I) Where, upon the death of an insured officer or key employee, the insurance proceeds are payable directly to the provider. In this case, the provider is a direct beneficiary. Insurance of this type is referred to as key-man insurance; and

(II) Where insurance on the lives of officers is voluntarily taken out as part of a mortgage loan agreement entered into for building construction and, upon the death of an insured officer, the proceeds are payable directly to the lending institution as a credit against the loan balance. In this case, the provider is an indirect beneficiary.

B. Types of insurance which are considered an allowable cost area—

(I) Where credit life insurance is required as part of a mortgage loan agreement. An example would be insurance on loans granted under certain federal programs; and

(II) Where the relative(s) or estate of the employee, excluding stockholders, partners and proprietors, is the beneficiary. [This type of insurance is considered to be] The MO HealthNet Division considers this type of insurance a fringe benefit and is an allowable cost area to the extent that the amount of coverage is reasonable.

2. Retirement plans.

A. Contributions to qualified retirement plans for the benefit of employees excluding stockholders, partners, and proprietors of the provider shall be allowable cost areas. [Interest] Facilities shall exclude interest income from funded pensions or retirement plans [shall be excluded] from consideration in determining the allowable cost area.

B. Amounts funded to pension and retirement plans, together with associated income, shall be recaptured if not actually paid when due, as an offset to expenses on the cost report form.

3. Deferred compensation plans.

A. Contributions for the benefit of employees, excluding stockholders, partners, and proprietors, under deferred compensation plans shall be all allowable cost areas when, and to the extent that, the costs are actually paid by the provider. Deferred compensation plans must be funded. Provider payments under unfunded deferred compensation plans will be considered as an allowable cost area only when paid to the participating employee and only to the extent considered reasonable.

B. Amounts paid by tax-exempt organizations to purchase tax-sheltered annuities for employees shall be treated as deferred compensation actually paid by the provider.

C. Amounts funded to deferred compensation plans, together with associated income [shall be recaptured] if not actually paid when due, as an offset to expenses on the cost report form.

(J) Education and Training Expenses.

1. The cost of on-the-job training [which] that directly benefits the quality of health care or administration at the facility shall be allowable. Off-the-job training involving extended periods exceeding five (5) continuous days is an allowable cost item only when specifically authorized in advance by the department.

2. Cost of education and training shall include incidental travel costs, but will not include leaves of absence or sabbaticals.

(K) Organizational Cost Items.

1. Organizational cost items may be included as an allowable cost area on an amortized basis.

2. Organizational cost items include the following: legal fees incurred in establishing the corporation or other organizations, necessary accounting fees, expenses of temporary directors, and organizational meetings of directors and stockholders, and fees paid to states of incorporation.

3. [Organizational costs shall be amortized] The provider shall amortize organizational costs ratably over a period of sixty (60) months beginning with the date of organization. When the provider enters the program more than sixty (60) months after the
date of organization, no organizational costs shall be recognized.

4. Where a provider did not capitalize organizational costs and has written off those costs in the year they were incurred, the provider cannot retroactively capitalize any part of these costs under the program.

5. Where a provider is organized within a five- (5-) year period prior to entering the program and has properly capitalized organizational costs using a sixty- (60-) month amortization period, no change in the rate of amortization is required. In this instance the unamortized portion of organizational costs is an allowable cost area under the program and shall be amortized over the remaining part of the sixty- (60-) month period.

6. For change in ownership after July 18, 1984, allowable amortization will be limited to the prior owner’s allowable unamortized portion of organizational cost.

(L) Advertising Costs. Advertising costs [which] that are reasonable, appropriate, and helpful in developing, maintaining, and furnishing services shall be an allowable cost area. The costs must be common and accepted occurrence in the field of the activity of the provider.

(M) Cost of Suppliers Involving Related Parties. Costs applicable to facilities, goods, and services furnished to a provider by a supplier related to the provider shall not exceed the lower of the cost to the supplier or the prices of comparable facilities, goods, or services obtained elsewhere. A provider shall identify suppliers related to it in the uniform cost report and the type-quantity and costs of facilities, goods, and services obtained from each supplier.

(N) Utilization Review. Incurred cost for the performance of required utilization review for ICF/IID is an allowable cost area. The expenditures must be for [the purpose of] providing utilization review on behalf of a Title XIX participant. [Utilization] The provider shall apportion utilization review costs incurred for Title XVIII and Title XIX [must be apportioned on the basis of] based on reimbursable participant days recorded for each program during the reporting period.

(O) Minimum Utilization. In the event the occupancy of a provider is below ninety percent (90%), the [provider] will calculate the following cost centers [will be calculated] as if the provider experienced ninety percent (90%) occupancy: laundry, housekeeping, general, administrative, and plant operation costs. In no case may the provider carry forward costs disallowed under this provision [be carried forward] to succeeding periods.

(P) Nonreimbursable Costs.

1. Bad debts, charity, and courtesy allowances are deductions from revenue and are not to be included as allowable costs.

2. Those services that are specifically provided by Medicare and MO HealthNet must be billed to those agencies.

3. Any costs incurred that are related to fund drives are not reimbursable.

4. Costs incurred for research purposes shall not be included as allowable costs.

5. The cost of services provided under the Title XX program, by contract or subcontract, is specifically excluded as an allowable item.

6. Attorney fees related to litigation involving state, local, or federal governmental entities and attorneys’ fees which are not related to the provision of LTC services, such as litigation related to disputes between or among owners, operators, or administrators.

7. Costs, such as legal fees, accounting and administration costs, travel costs, and the costs of feasibility studies, which are attributable to the negotiation or settlement of the sale or purchase of any capital asset by acquisition of merger for which any payment has been previously made under the program.

(Q) Other Revenues. Other revenues, including those listed that follow and excluding amounts collected under paragraph (5)(A)(8), will be deducted from the total allowable cost and must be shown separately in the cost report by use of a separate schedule if included in the gross revenue: income from telephone services; sale of employee and guest meals; sale of medical abstracts; sale of scrap and waste food or materials; rental income; cash, trade, quantity time, and other discounts; purchase rebates and refunds; recovery on insured loss; parking lot revenues; vending machine commissions or profit; sales from drugs to other than participants; income from investments of whatever type; and room reservation charges for temporary leave of absence days which are not covered services under section (5) of this rule. Failure by the provider to, in a readily ascertainable manner, separately account for any of the revenues specifically set out previously [in this rule in a readily ascertainable manner] in this rule, shall result in the provider’s termination from the program.

1. Interest income received from a funded depreciation account will not be deducted from allowable operating costs [provided that] if interest is applied to the replacement of the asset being depreciated.

2. Cost centers or operations specified by the provider in paragraph (7)(R)/3. (6)(R) of this rule shall not have their associated cost or revenues included in the covered costs or revenues of the facility.

3. Restricted and unrestricted funds.

A. “Restricted funds,” as used in this rule, mean those funds, cash or otherwise, including grants, gifts, taxes, and income from endowments, which the [provider] [must be used only] shall only use for a specific purpose designated by the donor. Those restricted funds [which] that are not transferred funds and are designated by the donor for paying operating costs will be offset from the total allowable expenses. If an administrative body has the authority to re-restrict restricted funds designated by the donor for paying operating costs, the [funds] provider will not [be] offset the funds from the total allowable expenses.

B. “Unrestricted funds,” as used in this rule, mean those funds, cash or otherwise, including grants, gifts, taxes, and income from endowments, that [are given] a donor gives to a provider without restriction [by the donor] as to their use. [These funds can be used] The provider can use these funds in any manner [desired by the provider]. However, those unrestricted funds [which] that are not transferred funds and [are used for paying] that the provider uses to pay operating costs will be offset from total allowable expenses.

C. Transferred funds as used in this rule are those funds appropriated through a legislative or governmental administrative body’s action, state or local, to a state or local government provider. The transfer can be state-to-state, state-to-local, or local-to-local provider. [These funds are not considered] The MO HealthNet Division does not consider these funds a grant or gift for reimbursement purposes, so [having] have no effect on the provider’s allowable cost under this plan.

(R) Apportionment of Costs to MO HealthNet Participant Residents.

1. [Provider’s] Providers shall apportion their allowable cost areas [shall be apportioned] between MO HealthNet program participant residents and other [patients] residents so that the share of allowable cost areas borne by the MO HealthNet program is based upon actual services received by MO HealthNet program participants.

2. To accomplish this apportionment, providers shall apply the ratio of [participant residents’ charges] patient days for MO HealthNet participants to the total patient [charges for the service of each ancillary department may be applied to the cost of this department. To this shall be added the cost of routine services for MO HealthNet program participant residents determined on the basis of a separate average cost per diem for general routine care areas or at the option of the provider on the basis of overall routine care area.

3. So that its charges may be allowable for use in apportioning costs under the program, each provider shall have an established charge structure which is applied uniformly to
each patient as services are furnished to the patient and which is reasonable and consistently related to the cost of providing these services/ days.

4.13. Average cost per diem for general routine services means the amount computed by dividing the total allowable patient costs for routine services by the total number of patient days of care rendered by the provider in the cost-reporting period.

5.14. A patient day of care is that period of service rendered a patient between the census-taking hours on two (2) consecutive days, including the twelve (12) temporary leave of absence days per any period of six (6) consecutive months as specifically covered under section (5) of this rule, the day of discharge being counted only when the patient was admitted the same day. [A census log shall be maintained] The provider shall maintain a census log in the facility for documentation purposes. Census shall be taken daily at mid-night. A day of care includes those overnight periods when a participant is away from the facility on a facility-sponsored group trip and remains under the supervision and care of facility personnel.

6.15. ICF/IID facilities that provide intermediate care services to MO HealthNet participants may establish distinct part cost centers in their facility provided that adequate accounting and statistical data required to separately determine the nursing care cost of each distinct part is maintained. Each distinct part may share the common services and facilities, such as management services, dietary, housekeeping, building maintenance, and laundry.

7.16. In no case may a provider’s allowable costs allocated to the MO HealthNet program include the cost of furnishing services to persons not covered under the MO HealthNet program.

(S) Return on Equity.
1. A return on a provider’s net equity shall be an allowable cost area.
2. The amount of return on a provider’s net equity shall not exceed twelve percent (12%) be calculated using the nursing home allowable percentage as defined in 13 CSR 70-10.015 Prospective Reimbursement Plan for Nursing Facility Services.
3. An owner’s net equity is comprised of investment capital and working capital. Investment capital includes the investment in building, property, and equipment (cost of land, mortgage payments toward principle, and equipment purchase less the accumulated depreciation). Working capital represents the amount of capital that is required to ensure proper operation of the facility.
4. The return on owner’s net equity shall be payable only to proprietary providers.
5. [A provider’s] The provider shall apportion its return on the owner’s net equity to the MO HealthNet program on the basis of, based on the provider’s MO HealthNet program reimbursable participant resident days of care to total resident days of care during the cost-reporting period. For the purpose of this calculation, total resident days of care shall be the greater of ninety percent (90%) of the provider’s certified bed capacity or actual occupancy during the cost year.

(T) Intermediate Care Facility for Individuals with Intellectual Disabilities Federal Reimbursement Allowance (ICF/IID FRA). The fee assessed to ICF/IID facilities in the state of Missouri for the privilege of doing business in the state will be an allowable cost.

18(7) Reporting Requirements.
A. Annual Cost Report.
1. Each provider shall establish a twelve- (12-) month fiscal period which is to be designated as the provider’s fiscal year. [An/ The provider shall submit an annual cost report for the fiscal year shall be submitted by the provider] to the department on forms to be furnished by the department for that purpose. [The] Each provider shall submit the completed cost report shall be submitted by the Department] by the first day of the sixth month following the close of the fiscal period.
2. Unless the provider has previously filed adequate and current documentation in the following areas [has been filed previously] with the department, authenticated copies of the following documents must be submitted by the provider with the cost reports: authenticated copies of all leases related to the activities of the facility; all management contracts, all contracts with consultants; federal and state income tax returns for the fiscal year; and documentation of expenditures, by line item, made under all restricted and unrestricted grants. For restricted grants, a statement verifying the restriction as specified by the donor.

3. [Adequate] The facility shall maintain adequate documentation for all line items on the uniform cost reports must be maintained by the facility] and must [be submitted] submit the document to the department upon request.
4. If a cost report is more than ten (10) days past due, payment shall may be withheld from the facility until the cost report is submitted. Upon receipt of a cost report prepared in accordance with this regulation, the department will release the withheld payments that were withheld will be released to the provider. For cost reports which are more than ninety (90) days past due, the department may terminate the provider’s MO HealthNet participation agreement and if terminated, retain all payments which have been withheld pursuant to this provision.

5. If a provider notifies, in writing, the director of the Institutional Reimbursement Unit of the division prior to the change of control, ownership, or termination of participation in the MO HealthNet program, the division will may withhold all remaining payments from the selling provider until the provider files the cost report is filed. The fully completed cost report with all required attachments and documentation is due the first day of the sixth month after the date of change of control, ownership, or termination. Upon receipt of a cost report prepared in accordance with this regulation, the department will release any withheld payment that was withheld will be released to the selling provider.

B. Certification of Cost Reports.
1. [The/ The facility must certify the] The facility must certify the accuracy and validity of any cost report must be certified]. Certification must be made by one (1) of the following persons (who must be authorized by the governing body of the facility to make the certification and will furnish proof of the authorization): an incorporated entity, an officer of the corporation; for a partnership, a partner; for a sole proprietorship or sole owner, the owner; or for a public facility, the chief administrative officer of the facility. The cost report must also be notarized by a licensed notary public.

2. Certification statement.
Form of Certification

Misrepresentation or falsifications of any information contained in this report may be punishable by fine, imprisonment, or both, under state or federal law.

Certification by officer or administrator of provider:
I hereby certify that I have read the above statement and that I have examined the accompanying cost report and supporting schedules prepared by

(Provider’s name(s) and number(s))

[for the cost report period beginning, ]

[20 and ending [ ]]

and that to the best of my knowledge and belief, it is a true, correct, and complete statement prepared from the books and records of the provider in accordance with applicable instructions, except as noted.

(Signature)     (Title)     (Date)

C. Adequacy of Records.
1. The provider must make available to the department or its duly authorized agent, including federal agents from Health and Human Services (HHS), at all reasonable times, the records as are necessary to permit review and audit of provider’s cost reports.
Failure to do so may lead to sanctions [stated in section (8) of this rule or other sanctions] available in section 70-3.030 or (9) of this rule.

2. [All] The provider shall retain all records associated with the preparation and documentation of the data associated with the cost report [must be retained] for seven (7) years from the cost report filing date.

(D) Accounting Basis.

2. Governmental institutions that operate on a cash or modified cash basis of accounting may continue to use those methods, provided [appropriate treatment of capital expenditures is made] the governmental institution treats capital expenditures appropriately.

(E) Audits.
1. [Cost reports shall be based] The provider shall base cost reports upon the provider's financial and statistical records [which] that must be capable of verification by audit.

2. If the provider has included the cost of a certified audit of the facility as an allowable cost item to the plan, a copy of that audit report and accompanying letter shall be submitted without deletions.

3. The annual cost report for the fiscal year of the provider may be subject to audit by the Department of Social Services or its contracted agents. Twelve- (12-) month cost reports for new construction facilities required to be submitted under section (4) of this rule may be audited by the department or its contracted agents prior to establishment of a permanent rate.

4. The department or authorized agent will conduct a desk review of all cost reports after submission by the provider and shall provide for on-site audits of facilities wherever their personnel note cost variances or exceptions [are noted by their personnel].

5. The department shall retain the annual cost report and any working papers relating to the audits of those cost reports for a period of not less than seven (7) full years from the date of submission of the report or completion of the audit.

6. Those providers having an annual Title XIX bed-day ratio on total bed days or certified beds of greater than sixty percent (60%) or an annual Title XIX payment of two hundred thousand dollars ($200,000) or more, or both, shall be required, for at least the first two (2) fiscal years of participation in the plan, to have an annual audit of their financial records by an independent certified public accountant. The auditor may issue a qualified audit report stating that confirmations of accounts receivable and accounts payable are not required by the plan. For the purposes of the paragraph, the Department of Social Services will only accept an unqualified opinion from a certified public accounting firm. A copy of the audit report must be submitted to the department to support the annual cost report of the facility.

(9)/(8) Sanctions and Overpayments.

(A) Sanctions may be imposed The department may impose sanctions against a provider in accordance with 13 CSR 70-3.030 and other federal or state statutes and regulations.

(B) In the case of overpayments to providers based on, but not limited to, field or audit findings or determinations based on a comprehensive operational review of the facility, the provider shall repay the overpayment in accordance with the provisions as set forth in 13 CSR 70-3.030.

(10)/(9) Exceptions.

(A) For those MO HealthNet-eligible participant-patients who have concurrent Medicare Part A skilled nursing facilities benefits available, MO HealthNet reimbursement for covered days of stay in a qualified facility will be based on the coinsurance as may be imposed under the Medicare Program.

(B) The Title XIX reimbursement rate for out-of-state providers shall be set by one (1) of the following methods:

1. For providers which provided prior authorized services of fewer than one thousand (1,000) patient days for Missouri Title XIX participants, the reimbursement rate shall be the rate paid for comparable services and level-of-care by the state in which the provider is located; and

2. For providers [which] that provide prior authorized services of one thousand (1,000) or more patient days for Missouri Title XIX participants, the reimbursement rate shall be the lower of—

A. The rate paid for comparable services and level-of-care by the state in which the provider is located; or

B. The rate calculated in [sections (4) and (6)] section (4) of this rule.

(10)/(10) Payment Assurance.

(A) The state will pay each provider, which furnished the services in accordance with the requirements of the state plan, the amount determined for services furnished by the provider according to the standards and methods set forth in these rules.

(B) Where third-party payment is involved, MO HealthNet will be the payor of last resort with the exception of state programs such as Vocational Rehabilitation and the Missouri Crippled Children’s Service. Procedures for remitting third-party payments are provided in the MO Healthnet program provider manuals.

(10)/(11) Provider Participation. Payments made in accordance with the standards and methods described in this rule are designed to enlist participation of a sufficient number of providers in the program so that eligible persons can receive medical care and services included in the state plan at least to the extent these services are available to the general public.

(10)/(12) Payment in Full. Participation in the program shall be limited to providers who accept as payment in full for covered services rendered to MO HealthNet participants, the amount paid in accordance with these rules and applicable copayments.

(10)/(13) Plan Evaluation. [Documentation will be maintained] The provider will maintain documentation to effectively monitor and evaluate experience during administration of this rule.

AUTHORITY: sections 208.153, 208.159, 208.201, and 660.017, RSMo 2016. For intervening history, please consult the Code of State Regulations. Emergency amendment filed Oct. 25, 2019, effective Nov. 8, 2019, expires May 5, 2020. A proposed amendment covering this same material is published in this issue of the Missouri Register.

PUBLIC COST: This emergency amendment will cost state agencies or political subdivisions approximately $1,023,030 in the time the emergency is effective.

PRIVATE COST: This proposed amendment will not cost private entities more than $500 in the time the emergency is effective.
FISCAL NOTE
PUBLIC COST

I. Department Title: Title 13 – Department of Social Services
Division Title: Division 70 – MO HealthNet Division
Chapter Title: Chapter 10 – Nursing Home Program

<table>
<thead>
<tr>
<th>Rule Number and Name:</th>
<th>13 CSR 70-10.030 Prospective Reimbursement Plan for Nonstate-Operated Facilities for ICF/IID Services</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type of Rulemaking:</td>
<td>Emergency Amendment</td>
</tr>
</tbody>
</table>

II. SUMMARY OF FISCAL IMPACT

<table>
<thead>
<tr>
<th>Affected Agency or Political Subdivision</th>
<th>Estimated Cost of Compliance in the Aggregate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Department of Mental Health</td>
<td>Cost for the time period the emergency amendment is effective is approximately $1,023,030</td>
</tr>
</tbody>
</table>

III. WORKSHEET

The annual cost of the rate rebase is approximately $818,421. The rate rebase is effective January 1, 2019; thus, the cost for the time period the emergency is effective is $1,023,030.

<table>
<thead>
<tr>
<th>Nonstate Operated ICF/IID(s)</th>
<th>Estimated Days</th>
<th>Est. Rate Increase/</th>
<th>Hold Harmless *</th>
<th>Estimated Impact</th>
</tr>
</thead>
<tbody>
<tr>
<td>Facility 1</td>
<td>3,172</td>
<td>$ 33.51</td>
<td>$ 106,294</td>
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</tr>
<tr>
<td>Facility 2</td>
<td>2,671</td>
<td>$ 69.87</td>
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<td>Facility 3</td>
<td>2,342</td>
<td>$ 78.08</td>
<td>$ 182,863</td>
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<td>Facility 4</td>
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<td>$ 29.98</td>
<td>$ 342,641</td>
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<tr>
<td>Facility 7</td>
<td>2,652</td>
<td>$ 0.00</td>
<td>$ 0.00</td>
<td></td>
</tr>
</tbody>
</table>

Total Annual Days / Cost Divided by 12 Months: 28,834 / 12 = $ 68,202

Months Paid in the Time the Emergency is Effective:
January 2019 - March 2020: 15

Cost in the Time the Emergency is Effective: $1,023,030

* Facilities that are “Hold Harmless” will not receive a rate increase but will continue to receive their current rate. See IV. Assumptions below for additional information.
IV. ASSUMPTIONS

The rebased rates are based on 2017 cost report data trended to 2019, the year that the rates become effective. A facility whose preliminary, recalculated rate is less than its current rate will continue to receive its current rate (i.e., Hold Harmless).

The estimated days are from the 2017 data. Since the nonstate-operated ICF/IID capacity has a stable census from year to year, the days from the 2017 base year do not require a utilization adjustment.
EMERGENCY AMENDMENT

22 CSR 10-2.020 General Membership Provisions. The Missouri Consolidated Health Care Plan is amending sections (3), (5), and (13).

PURPOSE: This amendment revises plan change criteria for Medicare Advantage Plan members, default enrollment procedures, clarifies disabled dependent eligibility, reporting of other health coverage, and renumbers as necessary.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting employees, retirees, officers, and their families enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to members as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register.

This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(3) Enrollment Procedures,

(A) Active Employee Coverage.

1. Statewide Employee Benefit Enrollment System (SEBES). A new employee must enroll or waive coverage through SEBES at www.sebes.mo.gov or through another designated enrollment system within thirty-one (31) days of his/her hire date or the date the employer notifies the employee that s/he is an eligible variable-hour employee. If enrolling a spouse or child(ren), proof of eligibility must be submitted as defined in section (5).

2. An active employee may elect, change, or cancel coverage for the next plan year during the annual open enrollment period that runs October 1 through October 31 of each year.

3. An active employee may elect or change coverage for himself/herself and/or for his/her spouse/child(ren) if one (1) of the following occurs:

   A. Occurrence of a life event, which includes marriage, birth, adoption, and placement of children. A special enrollment period of thirty-one (31) days shall be available beginning with the date of the life event. It is the employee’s responsibility to notify MCHCP of the life event.

   B. If paternity is necessary to establish the life event and was not established at birth, the date that paternity is established shall be the date of the life event; or

   C. Employer-sponsored group coverage loss. An employee or his/her spouse/child(ren) may enroll within sixty (60) days due to an involuntary loss of employer-sponsored coverage under one (1) of the following circumstances:

      (I) Employer-sponsored medical, dental, or vision plan terminates;

      (II) Eligibility for employer-sponsored coverage ends;

      (III) Employer contributions toward the premiums end; or

      (IV) COBRA coverage ends; or

   D. If an active employee or active employee’s spouse receives a court order stating s/he is responsible for covering a child, the active employee may enroll the child in an MCHCP plan within sixty (60) days of the court order.

4. Default enrollment.

   A. If an active employee is enrolled in the PPO [300 or/750, PPO 600/1250, or HSA Plan and does not complete enrollment during the open enrollment period, the employee and his/her dependents will be enrolled in the same plan enrolled in the prior year at the same level of coverage [in the PPO 1250 Plan provided through the vendor the employee is enrolled in, effective the first day of the next calendar year].

   B. If an active employee is enrolled in the Health Savings Account (HSA) Plan and does not complete enrollment during the open enrollment period, the employee and his/her dependents will be enrolled in the HSA Plan at the same level of coverage.

   C. If an active employee is enrolled in the TRICARE Supplemental Plan and does not complete enrollment during the open enrollment period, the employee and his/her dependents will be enrolled in the TRICARE Supplemental Plan at the same level of coverage.

   D. If a married state employee who is both MCHCP members who do not complete enrollment during the open enrollment period, will continue to meet one (1) family deductible and out-of-pocket maximum if they chose to do so during the previous plan year.

   E. If an active employee is enrolled in dental and/or vision coverage and does not complete open enrollment to cancel coverage or change the current level of coverage during the open enrollment period, the employee and his/her dependents will be enrolled at the same level of coverage in the same plan(s), effective the first day of the next calendar year.

5. If an active employee submits an Open Enrollment Worksheet or an Enroll/Change/Cancel form that is incomplete or contains obvious errors, MCHCP will notify the employee of such by mail, phone, or secure message. The employee must submit a corrected form to MCHCP by the date enrollment was originally due to MCHCP or ten (10) business days from the date the notice was mailed or sent by secure message or phone, whichever is later.

(B) Retiree Coverage.

1. To enroll or continue coverage for him/herself and his/her dependents or spouse/child(ren) at retirement, the employee must submit one (1) of the following:

   A. A completed enrollment form within thirty-one (31) days of retirement date even if the retiree is continuing coverage as a variable-hour employee after retirement. Coverage is effective on retirement date; or

   B. A completed enrollment form thirty-one (31) days before retirement date to have his/her first month’s retirement premium deducted and divided between his/her last two (2) payrolls and the option to pre-pay premiums through the cafeteria plan; or
C. A completed enrollment form within thirty-one (31) days of retirement date with proof of prior medical, dental, or vision coverage under a group or individual insurance policy for six (6) months immediately prior to his/her retirement if s/he chooses to enroll in an MCHCP Plan at retirement and has had insurance coverage for six (6) months immediately prior to his/her retirement.

2. A retiree may later add a spouse/child(ren) to his/her current coverage if one (1) of the following occurs:

A. Occurrence of a life event, which includes marriage, birth, adoption, and placement of children. A special enrollment period of thirty-one (31) days shall be available beginning with the date of the life event. It is the employee’s responsibility to notify MCHCP of the life event.

   (I) If paternity is necessary to establish the life event and was not established at birth, the date that paternity is established shall be the date of the life event; or
   
   B. Employer-sponsored group coverage loss. A retiree may enroll his/her spouse/child(ren) within sixty (60) days due to an involuntary loss of employer-sponsored coverage under one (1) of the following circumstances and the coverage was in place for twelve (12) months immediately prior to the loss:

   (I) Employer-sponsored medical, dental, or vision plan terminates;
   
   (II) Eligibility for employer-sponsored coverage ends;
   
   (III) Employer contributions toward the premiums end; or
   
   (IV) COBRA coverage ends.

3. If coverage was not maintained while on disability, the employee may enroll him/herself and his/her spouse/child(ren) within thirty-one (31) days of the date the employee is eligible for retirement benefits subject to the eligibility provisions herein.

4. A retiree may change from one (1) medical plan to another during open enrollment, but cannot add coverage for a spouse/child(ren). If a retiree is not already enrolled in medical, dental, and/or vision coverage, s/he cannot enroll in additional coverage during open enrollment.

5. A retiree enrolled in the Medicare Advantage Plan, may request to change to the PPO 750 Plan if the member is all of the following:

   A. A resident in a long term nursing facility;
   
   B. Eligible for Medicaid nursing home coverage, also known as “vendor coverage;” and
   
   C. Not a Qualified Medicare Beneficiary.


A. A retiree with Medicare and dependents with Medicare will be enrolled in the Medicare Advantage Plan.

   (I) If the retiree or a dependent becomes Medicare eligible in January of the next calendar year, they will be enrolled in the Medicare Advantage Plan.

   (II) If the retiree is not able to be enrolled in the Medicare Advantage Plan, [does not have Medicare Part B, and does not complete enrollment during the open enrollment period,] the retiree and his/her dependents without Medicare will be enrolled in the [PPO 1250 Plan provided through the vendor the retiree is enrolled in, effective the first day of the next calendar year] same plan enrolled in the prior year with the same level of coverage.

   [D. If a retiree without Medicare is enrolled in the HSA Plan and does not complete enrollment during the open enrollment period, the retiree and his/her dependents without Medicare will be enrolled in the HSA Plan through the vendor the retiree is enrolled in at the same level of coverage, effective the first day of the next calendar year.]

   [E./D. If a retiree without Medicare is currently enrolled in the TRICARE Supplemental Plan and does not complete enrollment during the open enrollment period, the retiree and his/her dependents will be enrolled in the TRICARE Supplemental Plan at the same level of coverage, effective the first day of the next calendar year.]

5/7. If a retiree is enrolled in dental and/or vision coverage and does not complete open enrollment during the open enrollment period, the retiree and his/her dependents will be enrolled at the same level of coverage in the same plan(s), effective the first day of the next calendar year.

7/8. If a retiree submits an Open Enrollment Worksheet, an Enroll/Change/Cancel form, or Retiree Enrollment form that is incomplete or contains obvious errors, MCHCP will notify the retiree of such by mail, phone, or secure message. The retiree must submit a corrected form to MCHCP by the date enrollment was originally due to MCHCP or ten (10) business days from the date the notice was mailed or sent by secure message or phone, whichever is later.

(C) Terminated Vested Coverage.

1. A terminated vested subscriber may later add a spouse/child(ren) to his/her coverage if one (1) of the following occurs:

   A. Occurrence of a life event, which includes marriage, birth, adoption, and placement of children. A special enrollment period of thirty-one (31) days shall be available beginning with the date of the life event. It is the employee’s responsibility to notify MCHCP of the life event.

   (I) If paternity is necessary to establish the life event and was not established at birth, the date that paternity is established shall be the date of the life event; or
   
   B. Employer-sponsored group coverage loss. A terminated vested subscriber may enroll his/her spouse/child(ren) within sixty (60) days due to an involuntary loss of employer-sponsored coverage under one (1) of the following circumstances and the coverage was in place for twelve (12) months immediately prior to the loss:

   (I) Employer-sponsored medical, dental, or vision plan terminates;
   
   (II) Eligibility for employer-sponsored coverage ends;
   
   (III) Employer contributions toward the premiums end; or
   
   (IV) COBRA coverage ends.

2. An enrolled terminated vested subscriber may change from one (1) medical plan to another during open enrollment but cannot add a spouse/child(ren) if the enrolled terminated vested subscriber is not already enrolled in medical, dental, and/or vision coverage, s/he cannot enroll in additional coverage during open enrollment.

3. A terminated vested member enrolled in the Medicare Advantage Plan, may request to change to the PPO 750 Plan if the member is all of the following:

   A. A resident in a long term nursing facility;
   
   B. Eligible for Medicaid nursing home coverage, also known as “vendor coverage;” and
   
   C. Not a Qualified Medicare Beneficiary.


A. A terminated vested subscriber with Medicare and dependents without Medicare will be enrolled in the Medicare Advantage Plan.
(I) If the terminated vested subscriber or a dependent becomes Medicare eligible in January of the next calendar year, they will be enrolled in the Medicare Advantage Plan.

(II) If the terminated vested subscriber [does not have Medicare Part B, and does not complete enrollment during the open enrollment period] is not able to be enrolled in the Medicare Advantage Plan, the terminated vested subscriber and his/her dependents without Medicare will be enrolled in the [PPO 1250 plan provided through the vendor the terminated vested subscriber is enrolled in, effective the first day of the next calendar year] same plan enrolled in the prior year with the same level of coverage.

B. If a terminated vested subscriber without Medicare is enrolled in the PPO [300] 750, [or] PPO [600] 1250, or HSA Plan and does not complete enrollment during the open enrollment period, the terminated vested subscriber and his/her dependents without Medicare will be enrolled in the [PPO 1250 Plan provided through the vendor the terminated vested subscriber is enrolled in, effective the first day of the next calendar year] same plan enrolled in the prior year with the same level of coverage.

C. If a terminated vested subscriber without Medicare is enrolled in the HSA Plan and does not complete enrollment during the open enrollment period, the terminated vested subscriber and his/her dependents without Medicare will be enrolled in the HSA Plan through the vendor the terminated vested subscriber is enrolled in effective the first day of the next calendar year, at the same level of coverage.

D. If a terminated vested subscriber without Medicare is enrolled in the TRICARE Supplemental Plan and does not complete enrollment during the open enrollment period, the terminated vested subscriber and his/her dependents without Medicare will be enrolled in the TRICARE Supplemental Plan effective the first day of the next calendar year, at the same level of coverage.

E. If a terminated vested subscriber is enrolled in dental and/or vision coverage and does not complete open enrollment during the open enrollment period, the employee and his/her dependents will be enrolled at the same level of coverage in the same plan(s), effective the first day of the next calendar year.

F. If a terminated vested subscriber submits an Open Enrollment Worksheet, an Enroll/Change/Cancel form, or Terminated Vested Enrollment form that is incomplete or contains obvious errors, MCHCP will notify the terminated vested subscriber of such by mail, phone, or secure message. The terminated vested subscriber must submit a corrected form to MCHCP by the due date enrollment was originally due to MCHCP or ten (10) business days from the date the notice was mailed or sent by secure message or phone, whichever is later.

D. Long-Term Disability Coverage.

1. A long-term disability subscriber may add a spouse/child(ren) to his/her current coverage if one (1) of the following occurs:

A. Occurrence of a life event, which includes marriage, birth, adoption, and placement of children. A special enrollment period of thirty-one (31) days shall be available beginning with the date of the life event. It is the employee’s responsibility to notify MCHCP of the life event.

B. Employer-sponsored group coverage loss. A long-term disability subscriber may enroll his/her spouse/child(ren) within sixty (60) days due to an involuntary loss of employer-sponsored coverage under one (1) of the following circumstances and the coverage was in place for twelve (12) months immediately prior to the loss:

   I. Employer-sponsored medical, dental, or vision plan terminates;

   II. Eligibility for employer-sponsored coverage ends;

   III. Employer contributions toward the premiums end; or

   IV. COBRA coverage ends.

2. An enrolled long-term disability subscriber may change from one (1) medical plan to another during open enrollment but cannot add a spouse/child(ren). If an enrolled long-term disability subscriber is not already enrolled in medical, dental, and/or vision coverage, s/he cannot enroll in additional coverage during open enrollment.

3. A long-term disability member enrolled in the Medicare Advantage Plan, may request to change to the PPO 750 Plan if the member is all of the following:

A. A resident in a long term nursing facility;

B. Eligible for Medicaid nursing home coverage, also known as “vendor coverage;” and

C. Not a Qualified Medicare Beneficiary.

4. Default enrollment.

A. A long-term disability subscriber with Medicare and dependents with Medicare will be enrolled in the Medicare Advantage Plan.

B. If a long-term disability subscriber without Medicare is enrolled in the PPO [300] 750, [or] PPO [600] 1250, or HSA Plan and does not complete enrollment during the open enrollment period, the long-term disability subscriber and his/her dependents without Medicare will be enrolled in the [PPO 1250 Plan provided through the vendor the long-term disability subscriber is enrolled in, effective the first day of the next calendar year] same plan enrolled in the prior year with the same level of coverage.

C. If a long-term disability subscriber with Medicare has a non-Medicare dependent enrolled in the PPO [300] 750, [or] PPO [600] 1250, or HSA Plan and does not complete enrollment during the open enrollment period, [and has dependents who are not covered by Medicare], the long-term disability subscriber and his/her dependents without Medicare will be enrolled in the [PPO 1250 Plan provided through the vendor the long-term disability subscriber is enrolled in, effective the first day of the next calendar year] same plan enrolled in the prior year with the same level of coverage.

5. Eligibility for employer-sponsored coverage ends.
Emergency Rules

Enrollment Worksheet or an Enroll/Change/Cancel form that is incomplete or contains obvious errors, MCHCP will notify the long-term disability subscriber of such by mail, phone, or secure message. The long-term disability subscriber must submit a corrected form to MCHCP by the date enrollment was originally due to MCHCP or ten (10) business days from the date the notice was mailed or sent by secure message or phone, whichever is later.

(E) Survivor Coverage.

1. A survivor without Medicare must submit a survivor enrollment form [and a copy of the death certificate] within thirty-one (31) days of the first day of the month after the death of the employee. The survivor must not elect coverage within thirty-one (31) days of the first day of the month after the death of the employee, s/he cannot enroll at a later date.

2. A survivor with Medicare will be automatically enrolled as a survivor following the death of the employee.

(A) If the survivor does not elect coverage within thirty-one (31) days of the first day of the month after the death of the employee, s/he cannot enroll at a later date.

(B) If the survivor marries, has a child, adopts a child, or a child is placed with the survivor, the spouse/child(ren) must be added within thirty-one (31) days of birth, adoption, placement, or marriage.

(C) If eligible spouse/child(ren) are not enrolled when first eligible, they cannot be enrolled at a later date.

3. A survivor may later add a spouse/child(ren) to his/her current coverage if one (1) of the following occurs:

(I) Occurrence of a life event, which includes marriage, birth, adoption, and placement of children. A special enrollment period of thirty-one (31) days shall be available beginning with the date of the life event. It is the employee’s responsibility to notify MCHCP of the life event.

(II) If paternity is necessary to establish the life event and was not established at birth, the date that paternity is established shall be the date of the life event; or

(B) Eligible for Medicaid nursing home coverage. A survivor may enroll his/her spouse/child(ren) within sixty (60) days due to an involuntary loss of employer-sponsored coverage under one (1) of the following circumstances and the coverage was in place for twelve (12) months immediately prior to the loss:

(I) Employer-sponsored medical, dental, or vision plan terminates;

(II) Eligibility for employer-sponsored coverage ends;

(III) Employer contributions toward the premiums end; or

(IV) COBRA coverage ends.

(C) If a survivor without Medicare is enrolled in the PPO 750, [or] PPO 600/1250, or HSA Plan and does not complete enrollment during the open enrollment period, the survivor and his/her dependents without Medicare will be enrolled in the [PPO 1250 Plan provided through the vendor the survivor is enrolled in, effective the first day of the next calendar year] same plan enrolled in the prior year with the same level of coverage.

1. If the survivor or a dependent becomes Medicare eligible in January of the next calendar year, they will be enrolled in the Medicare Advantage Plan.

2. If the survivor [does not have Medicare Part B, and does not complete enrollment during the open enrollment period] is not able to be enrolled in the Medicare Advantage Plan, the survivor and his/her dependents without Medicare will be enrolled in the [PPO 1250 Plan provided through the vendor the retiree is enrolled in, effective the first day of the next calendar year] same plan enrolled in the prior year with the same level of coverage.

3. If a survivor with Medicare has a non-Medicare dependent [is] enrolled in the [PPO 300/750, [or] PPO 600/1250, or HSA Plan and does not complete enrollment during the open enrollment period [and has dependents who are not covered by Medicare], the survivor and his/her dependents without Medicare will be enrolled in the [PPO 1250 Plan provided through the vendor the retiree is enrolled in, effective the first day of the next calendar year] same plan enrolled in the prior year with the same level of coverage.

4. A resident in a long term nursing facility;

5. If a survivor without Medicare is enrolled in the [HSA Plan and does not complete enrollment during the open enrollment period, the survivor and his/her dependents without Medicare will be enrolled in the [PPO 1250 Plan provided through the vendor the survivor is enrolled in, effective the first day of the next calendar year] same plan enrolled in the prior year with the same level of coverage.

6. A resident in a long term nursing facility;

7. Not a Qualified Medicare Beneficiary.

/F/.. If a survivor is enrolled in dental and/or vision coverage and does not complete open enrollment during the open enrollment period, the survivor and his/her dependents will be enrolled at the same level of coverage in the same plan(s), effective the first day of the next calendar year.

/G/.. The employee may enroll his/her permanently disabled child when first eligible or an enrolled permanently disabled dependent turning age twenty-six (26) years and may continue coverage beyond age twenty-six (26) years, provided the following documentation is submitted to the plan prior to the end of the month the
dependent’s twenty-sixth birthday for the enrolled permanently disabled dependent or within thirty-one (31) days of enrollment of [a new employee and his/her] the permanently disabled child.

A. Evidence from the Social Security Administration (SSA) that the permanently disabled dependent or child was entitled to and receiving disability benefits prior to turning age twenty-six (26) years; and

B. A benefit verification letter dated within the last twelve (12) months from the SSA confirming the child is still considered disabled.

2. If a disabled dependent or child over the age of twenty-six (26) years is determined to be no longer disabled by the SSA, coverage will terminate the last day of the month in which the disability ends or will never take effect for new enrollment requests.

3. Once the disabled dependent’s coverage is cancelled or terminated, s/he will not be able to enroll at a later date.

(13) Members are required to disclose to the claims administrator whether or not they have other health coverage and, if so, information about the coverage. [A member may submit this information to the claims administrator by phone, fax, mail, or online. Dependent claims will be denied if the disclosure is not made.] Once the information is received, claims will be reprocessed subject to all applicable rules.


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 2—State Membership

EMERGENCY RESCISSION

22 CSR 10-2.045 Plan Utilization Review Policy. This rule established the policy of the board of trustees in regard to the Plan Utilization Review Policy of the Missouri Consolidated Health Care Plan.

PURPOSE: This rule is being rescinded and readopted to reflect changes due to a new third party administrator.

EMERGENCY STATEMENT: This emergency rescission must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency rescission is necessary to serve a compelling governmental interest of protecting members (employees, retirees, officers, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this rescission be filed as an emergency rescission to maintain the integrity of the current health care plan. This emergency rescission fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to members as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency rescission reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed rescission, which covers the same material, is published in this issue of the Missouri Register. This emergency rescission complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency rescission was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.


PUBLIC COST: This emergency rescission will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency rescission will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 2—State Membership

EMERGENCY RULE

22 CSR 10-2.045 Plan Utilization Review Policy

PURPOSE: This rule establishes the policy of the board of trustees in regard to the Plan Utilization Review Policy of the Missouri Consolidated Health Care Plan Medical Plans.

EMERGENCY STATEMENT: This emergency rule must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency rule is necessary to serve a compelling governmental interest of protecting members (employees, retirees, officers, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this rule be filed as an emergency rule to maintain the integrity of the current health care plan. This emergency rule fulfills the compelling
governmental interest of offering access to more convenient and affordable medical services to members as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency rule reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed rule, which covers the same material, is published in this issue of the Missouri Register. This emergency rule complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency rule was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(1) Clinical Management—Certain benefits are subject to a utilization review (UR) program. The program has the following components:

(A) Preauthorization—The claims administrator must authorize some services in advance. Preauthorization is to determine if the procedure or treatment is medically necessary. The claims administrator will determine what procedures or treatments are subject to preauthorization. Without preauthorization, any claim that requires preauthorization will be denied for payment. Members who have another primary carrier, or who are enrolled in the Medicare Advantage Plan are not subject to this provision except for those services that are not covered by the other primary carrier, but are otherwise subject to preauthorization under this rule. Preauthorizations found to have a material misrepresentation or intentional or negligent omission about the person’s health condition or the cause of the condition may be rescinded.

1. A list of medical services for which preauthorization is required may be obtained at any time from the claims administrator.

2. The following pharmacy services included in the prescription drug plan for non-Medicare primary members are subject to preauthorization:

A. Second-step therapy medications that skip the first-step medication trial;
B. Specialty medications;
C. Medications that may be prescribed for several conditions, including some for which treatment is not medically necessary;
D. Medication refill requests that are before the time allowed for refill;
E. Medications that exceed drug quantity and day supply limitations; and
F. Medications with costs exceeding nine thousand nine hundred ninety-nine dollars and ninety-nine cents ($9,999.99) at retail or the mail order pharmacy and one hundred forty-nine dollars and ninety-nine cents ($149.99) for compound medications at retail or the mail order pharmacy.

3. Preauthorization timeframes.

A. A benefit determination for non-urgent preauthorization requests will be made within thirty-six (36) hours, which will include one (1) business day of the receipt of the request. If the information necessary to make a benefit determination is not received, the claims administrator will notify the member and provider of any necessary extension. The provider will be given forty-five (45) calendar days from receipt of the extension notice to respond with additional information. Once the information is received or the forty-five (45) days have elapsed, a determination will be made within thirty-six (36) hours which will include one (1) business day.

B. A benefit determination for urgent preauthorization requests will be made as soon as possible based on the clinical situation, but in no case later than one (1) business day of the receipt of all necessary information;

(B) Concurrent Review—The claims administrator will monitor the medical necessity of an inpatient admission to certify the necessity of the continued stay in the hospital. Members who have another primary carrier, including Medicare, are not subject to this provision;  

(C) Retrospective Review—Reviews to determine coverage after services have been provided to a member. The retrospective review is not limited to an evaluation of medical necessity, reimbursement levels, accuracy and adequacy of documentation or coding, or settling of payment. The claim administrator shall have the authority to correct payment errors when identified under retrospective review;

(D) Pre-determination—Determination of coverage by the claims administrator prior to services being provided. A provider may voluntarily request a pre-determination. A pre-determination informs the provider of whether, and under which circumstances, a procedure or service is generally a covered benefit under the plan. A pre-determination that a procedure or service may be covered under the plan does not guarantee payment; and

(E) Case Management—A voluntary process to assess, coordinate, and evaluate options and services of members with catastrophic and complex illnesses. A case manager will help members understand what to expect during the course of treatment, help establish collaborative goals, complete assessments to determine needs, interface with providers, and negotiate care. Members are identified for case management through claim information, length of hospital stay, or by referral. The case manager will dismiss the member from case management once the case manager determines that objectives have been met.


PUBLIC COST: This emergency rule will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency rule will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—Missouri Consolidated Health Care Plan
Division 10—Health Care Plan
Chapter 2—State Membership

Emergency Amendment

22 CSR 10-2.046 PPO 750 Plan Benefit Provisions and Covered Charges. The Missouri Consolidated Health Care Plan is amending sections (5), (11), (12), and (14).

PURPOSE: This amendment adds one hundred percent (100%) coverage for sterilization procedure for men, revises maximum plan payment, timely filing time frame, and services performed in another country for the PPO 750 Plan.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (employees, retirees, officers, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help
Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
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EMERGENCY AMENDMENT

22 CSR 10-2.047 PPO 1250 Plan Benefit Provisions and Covered Charges. The Missouri Consolidated Health Care Plan is amending sections (5), (11), (12), and (14).

PURPOSE: This amendment adds one hundred percent (100%) coverage for sterilization procedure for men, revises maximum plan payment, timely filing timeframe, and services performed in another country for the PPO 1250 Plan.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (employees, retirees, officers, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to members as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(5) The following services are not subject to deductible, coinsurance, or copayment requirements and will be paid at one hundred percent (100%) when provided by a network provider:

(A) A newborn’s initial hospitalization until discharge or transfer to another facility if the mother is a Missouri Consolidated Health Care Plan (MCHCP) member at the time of birth; [and]

(D) Four (4) Diabetes Self-Management Education visits[.]; and

(E) Sterilization procedure for men.

(11) Maximum plan payment—non-network medical claims that are not otherwise subject to a contractual discount arrangement are processed at one hundred ten percent (110%) of Medicare reimbursement for non-network professional claims and following the claim administrator’s standard practice for non-network facility claims. Members may be held liable for the amount of the fee above the allowed amount.

(12) Any claim must be initially submitted within twelve (12) months following the date of service, unless other specified in the network provider contract. The plan reserves the right to deny claims not timely filed. A provider initiated correction to the originally filed claim must be submitted within the timeframe agreed in the provider contract, but not to exceed three hundred sixty-five (365) days from adjudication of the originally filed claim. Any claims reprocessed as primary based on action taken by Medicare or Medicaid must be initiated within three (3) years of the claim being incurred.

(14) Services performed in a country other than the United States may be covered if the service is included in 22 CSR 10-2.055. Emergency and urgent care services are covered as a network benefit. All other non-emergency services are covered as [a non-network benefit] determined by the claims administrator. If the service is provided by a non-network provider, the member may be required to provide payment to the provider and then file a claim for reimbursement subject to timely filing limits.


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.
timely filed. A provider initiated correction to the originally filed claim must be submitted within the timeframe agreed in the provider contract, but not to exceed three hundred sixty-five (365) days from adjudication of the originally filed claim. Any claims reprocessed as primary based on action taken by Medicare or Medicaid must be initiated within three (3) years of the claim being incurred.

(14) Services performed in a country other than the United States may be covered if the service is included in 22 CSR 10-2.055. Emergency and urgent care services are covered as a network benefit. All other non-emergency services are covered as [a non-network benefit] determined by the claims administrator. If the service is provided by a non-network provider, the member may be required to provide payment to the provider and then file a claim for reimbursement subject to timely filing limits.


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
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EMERGENCY AMENDMENT

22 CSR 10-2.053 Health Savings Account Plan Benefit Provisions and Covered Charges. The Missouri Consolidated Health Care Plan is amending sections (3), (14), (15), (17), and adding section (9).

PURPOSE: This amendment revises the Health Savings Account (HSA) Plan individual family member out-of-pocket maximum, adds one hundred percent (100%) coverage after deductible is met for sterilization procedure for men, revises maximum plan payment, timely filing timeframe, services performed in another country, and remuners as necessary.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (employees, retirees, officers, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to members as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(3) Out-of-pocket maximum.

(A) The family out-of-pocket maximum applies when two (2) or more family members are covered. The family out-of-pocket maximum must be met before the plan begins to pay one hundred percent (100%) of all covered charges for any covered family member. Out-of-pocket maximums are per calendar year, as follows:

1. Network out-of-pocket maximum for individual—four thousand nine hundred fifty dollars ($4,950);
2. Network out-of-pocket maximum for family—nine thousand nine hundred dollars ($9,900). Any individual family member need only incur a maximum of [seven thousand nine hundred dollars ($7,900)] eight thousand one hundred fifty dollars ($8,150) before the plan begins paying one hundred percent (100%) of covered charges for that individual;
3. Non-network out-of-pocket maximum for individual—nine thousand nine hundred dollars ($9,900); and

(9) Sterilization procedure for men is paid at one hundred percent (100%) when provided by a network provider after deductible is met.

(/9)/(10) Newborn’s claims will be subject to deductible and coinsurance.

(/10)(/11) Married, active employees who are MCHCP subscribers and have enrolled children may meet only one (1) family deductible and out-of-pocket maximum. Both spouses must enroll in the same medical plan option through the same carrier, and each must provide the other spouse’s Social Security number (SSN) and report the other spouse as eligible for coverage when newly hired and during the open enrollment process. In the medical plan vendor and pharmacy benefit manager system, the spouse with children enrolled will be considered the subscriber and the spouse that does not have children enrolled will be considered a dependent. If both spouses have children enrolled the spouse with the higher Social Security number (SSN) will be considered the subscriber. Failure to report an active employee spouse when newly hired and/or during open enrollment will result in a separate deductible and out-of-pocket maximum for both active employees.

(/11)(/12) Each subscriber will have access to payment information of the family unit only when authorization is granted by the adult covered dependent(s).

(/12)(/13) Expenses toward the deductible and out-of-pocket maximum will be transferred if the member changes non-Medicare medical plans or continues enrollment under another subscriber’s non-Medicare medical plan within the same plan year.

(/13)(/14) Maximum plan payment—Non-network medical claims that are not otherwise subject to a contractual discount arrangement are processed at one hundred ten percent (110%) of Medicare reimbursement for non-network professional claims and following the
Any claim must be initially submitted within twelve (12) months following the date of service, unless other specified in the network provider contract. The plan reserves the right to deny claims not timely filed. A provider initiated correction to the originally filed claim must be submitted within the timeframe agreed in the provider contract, but not to exceed three hundred sixty-five (365) days from adjudication of the originally filed claim. Any claims reprocessed as primary based on action taken by Medicare or Medicaid must be initiated within three (3) years of the claim being incurred.

For a member who is an inpatient on the last calendar day of a plan year and remains an inpatient into the next plan year, the prior plan year’s applicable deductible and/or coinsurance amounts will apply to the in-hospital facility and related ancillary charges until the member is discharged.

Services performed in a country other than the United States may be covered if the service is included in 22 CSR 10-2.055. All other non-emergency services are covered as a network benefit. Emergency and urgent care services are covered as a network benefit. States may be covered if the service is included in 22 CSR 10-2.055.

For a member who is an inpatient on the last calendar day of a plan year and remains an inpatient into the next plan year, the prior plan year’s applicable deductible and/or coinsurance amounts will apply to the in-hospital facility and related ancillary charges until the member is discharged.


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 2—State Membership

EMERGENCY AMENDMENT

22 CSR 10-2.055 Medical Plan Benefit Provisions and Covered Charges. The Missouri Consolidated Health Care Plan is amending sections (2) and (3).
PURPOSE: This amendment revises the medical plan benefits for transition of care, allergy testing and immunotherapy, bariatric surgery, bone growth stimulators, cardiac rehabilitation, chelation therapy, chiropractic rehabilitation, cochlear implant, denial care, emergency room services, genetic counseling, hearing aids, hospice care and palliative services, hospital, injections, maternity coverage, nutrition therapy, orthognathic or jaw surgery, orthotics, and reminders as necessary.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (employees, retirees, officers, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to members as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

I(2) Transition of Care. A transition of care option is available for members using a hospital or dialysis facility that loses network status during the plan year. A subscriber and his/her dependents using a hospital or dialysis facility that loses network status during the plan year may apply for a ninety- (90-) day transition of care to continue receiving network benefits with that hospital or dialysis facility. The request for consideration must be submitted to the medical plan within forty-five (45) days of the last day the hospital or dialysis facility was a contracted network provider, to be eligible for transition of care benefits. A subscriber and his/her dependents may apply for additional days beyond the ninety- (90-) day transition if care is related to a moderate or high risk pregnancy, if care is during a member’s second or third trimester of pregnancy, or up to eight (8) weeks postpartum. The subscriber and his/her dependents must apply for additional transition of care days prior to the end of the initial ninety- (90-) day transition of care period. Most routine services, treatment for stable conditions, minor illnesses, and elective surgeries will not be covered by transition of care benefits. The rate of payment during the transitional period shall be the fee paid prior to leaving the network. Benefits eligible for transition of care include:

(A) Upcoming surgery or prospective transplant;
(B) Services for women in their second or third trimester of pregnancy or up to eight (8) weeks postpartum;
(C) Services for women who have been diagnosed as potentially having a moderate- or high-risk pregnancy;
(D) Home nursing care;
(E) Radiation therapy;
(F) Dialysis;
(G) Durable medical equipment;
(H) Cancer treatment;
(I) Clinical trials;
(J) Physical, speech, or occupational therapy;
(K) Hospice care;
(L) Bariatric surgery, and follow-up per criteria covered under the plan;
(M) Inpatient hospitalization at the time of the network change;
(N) Mental health services; or
(O) Related follow-up services within three (3) months of an acute injury or surgery.

(2) Transition of Care. A transition of care option is available for members who seek to continue to remain under the care of an non-network provider who was treating them prior to the provider losing network status. A subscriber and his/her dependents may request to continue receiving care at the network benefit level. If approved, the member will be eligible to continue care with the current non-network provider at the network benefit level for a period of time until it is medically appropriate for the member to transfer care to a network provider. The rate of payment during the transitional period shall be the fee paid prior to leaving the network. The following benefits are eligible for transition of care as determined by the claims administrator:

(A) Upcoming surgery or prospective transplant;
(B) Services for women in their third trimester of pregnancy;
(C) Radiation therapy;
(D) Dialysis;
(E) Cancer treatment;
(F) Physical, speech, or occupational therapy;
(G) Hospice care;
(H) Inpatient hospitalization at the time of the network change;

(3) Covered Charges Applicable to the PPO 750 Plan, PPO 1250, and HSA Plan.

(i) Services in a country other than the United States. Emergency room and urgent care medical services are covered at the network benefit. All other non-emergency services are covered at the non-network benefit.

(ii) Plan benefits for the PPO 750 Plan, PPO 1250, and HSA Plan are as follows:

1. Allergy Testing and Immunotherapy. Allergy testing and allergy immunotherapy are considered medically necessary for members with clinically significant allergic symptoms. The following tests and treatments are covered:

[A. Epicutaneous (scratch, prick, or puncture) when Immunoglobulin E (IgE) mediated reactions occur to any of the following:
(i) Foods;
(ii) Hymenoptera venom (stinging insects);
(iii) Inhalants; or
(iv) Specific drugs (penicillins and macromolecular agents);

B. Intradermal (Intracutaneous) when IgE-mediated reactions occur to any of the following:
(i) Foods;
(ii) Hymenoptera venom (stinging insects);]
(I) Ingestion (Oral) Challenge Test for any of the following:
(i) Food or other substances; or
(ii) Drugs when all of the following are met:
   (a) History of allergy to a particular drug;
   (b) There is no effective alternative drug; and
   (c) Treatment with that drug class is essential;

J. In Vitro IgE Antibody Tests (RAST, MAST, FAST, ELISA, immunoCAP) are covered for any of the following:
(i) Allergic broncho-pulmonary aspergillosis (ABPA) and certain parasitic diseases;
(ii) Food allergy;
(iii) Hymenoptera venom allergy (stinging insects);
(iv) Inhalant allergy; or
(v) Specific drugs;
K. Total Serum IgE for diagnostic evaluation in members with known or suspected ABPA and/or hyper IgE syndrome;
L. Lymphocyte transformation tests such as lymphocyte mitogen response test, PHE stimulation test, or lymphocyte antigen response assay are covered for evaluation of persons with any of the following suspected conditions:
(i) Sensitivity to beryllium;
(ii) Congenital or acquired immunodeficiency diseases affecting cell-mediated immunity, such as severe combined immunodeficiency, common variable immunodeficiency, X-linked immunodeficiency with hyper IgM, Nijmegen breakage syndrome, reticular dysgenesis, DiGeorge syndrome, Nezelof syndrome, Wiscott-Aldrich syndrome, ataxia telangiectasia, and chronic mucocutaneous candidiasis;
(iii) Thymoma; and
(iv) To predict allograft compatibility in the transplant setting;
M. Allergy retesting: routine allergy retesting is not considered medically necessary;

N. Allergy immunotherapy is covered for the treatment of any of the following IgE-mediated allergies:
(i) Allergic (extrinsic) asthma;
(ii) Dust mite atopic dermatitis;
(iii) Hymenoptera (bees, hornets, wasps, fire ants) hypersensitivity (hymenoptera); or

O. Other treatments: the following other treatments are covered:
(i) Rapid, rush, cluster, or acute desensitization for members with any of the following conditions:
   (a) IgE antibodies to a particular drug that cannot be treated effectively with alternative medications;
   (b) Insect sting (e.g., wasps, hornets, bees, fire ants) hypersensitivity (hymenoptera); or
   (c) Members with moderate to severe allergic rhinitis who need treatment during or immediately before the season of the affecting allergy;

(ii) Rapid desensitization is considered experimental and investigational for other indications;

P. Epinephrine kits, to prevent anaphylactic shock for members who have had life-threatening reactions to insect stings, foods, drugs, or other allergens; have severe asthma or if needed during immunotherapy;

2. Ambulance service. The following ambulance transport services are covered:
A. By ground to the nearest appropriate facility when other means of transportation would be contraindicated;
B. By air to the nearest appropriate facility when the member's medical condition is such that transportation by either basic or advanced life support ground ambulance is not appropriate or contraindicated;

3. Applied Behavior Analysis (ABA) for Autism;

4. Bariatric surgery. Bariatric surgery is covered when all of the following requirements have been met:

   [A. The surgery is performed at a facility accredited by the Metabolic and Bariatric Surgery Accreditation and Quality Improvement Program (MBSAQIP) for the billed procedure;]

   B. The following open or laparoscopic bariatric surgery procedures are covered:
   (i) Roux-en-Y gastric bypass;
   (ii) Sleeve gastrectomy;
   (iii) Biliopancreatic diversion with duodenal switch for individuals with a body mass index (BMI) greater than fifty (50);

   (IV) Adjustable silicone gastric banding and adjustments of a silicone gastric banding to control the rate of weight loss and/or treat symptoms secondary to gastric restriction following an adjustable silicone gastric banding procedure;

   (V) Surgical reversal of bariatric surgery when complications of the original surgery (e.g., stricture, pouch dilatation, erosion, or band slippage) cause abdominal pain, inability to eat or drink, or cause vomiting of prescribed meals;

   (VI) Revision of a previous bariatric surgical procedure or conversion to another procedure due to inadequate weight loss when one (1) of the following specific criteria has been met:
   (a) There is evidence of full compliance with the previously prescribed post-operative dietary and exercise program; or
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(b) There is documented clinical testing demonstrating technical failure of the original bariatric surgical procedure which caused the individual to fail achieving adequate weight loss of at least fifty percent (50%) of excess body weight or failure to achieve body weight to within thirty percent (30%) of ideal body weight at least two (2) years following the original surgery;

C. All of the following criteria have been met:
(I) The member is eighteen (18) years or older or has reached full skeletal growth, and has evidence of one (1) of the following:
(a) BMI greater than forty (40); or
(b) BMI between thirty-five (35) and thirty-nine and nine tenths (39.9) and one (1) or more of the following:
   I. Type II diabetes;
   II. Cardiovascular disease such as stroke, myocardial infarction, stable or unstable angina pectoris, hypertension, or coronary artery bypass; or
   III. Life-threatening cardiopulmonary problems such as severe sleep apnea, Pickwickian syndrome, or obesity-related cardiomyopathy; and

(II) Demonstration that dietary attempts at weight control have been ineffective through completion of a structured diet program. Commercial weight loss programs are acceptable if completed under the direction of a provider or registered dietitian and documentation of participation is available for review. One (1) structured diet program for six (6) consecutive months or two (2) structured diet programs for three (3) consecutive months each within a two- (2-) year period prior to the request for the surgical treatment of morbid obesity are sufficient. Provider-supervised programs consisting exclusively of pharmacological management are not sufficient; and

(III) A thorough multidisciplinary evaluation within the previous twelve (12) months, which include all of the following:
   (a) An evaluation by a bariatric surgeon recommending surgical treatment, including a description of the proposed procedure and all of the associated current procedural terminology codes;
   (b) A separate medical evaluation from a provider other than the surgeon recommending surgery that includes a medical clearance for bariatric surgery;
   (c) Completion of a psychological examination from a mental health provider evaluating the member’s readiness and fitness for surgery and the necessary post-operative lifestyle changes. After the evaluation, the mental health provider must provide clearance for bariatric surgery; and
   (d) A nutritional evaluation by a provider or registered dietitian;

5. Blood storage. Storage of whole blood, blood plasma, and blood products is covered in conjunction with medical treatment that requires immediate blood transfusion support;

6. Bone Growth Stimulators. Implantable bone growth stimulators are covered as an outpatient surgery benefit. The following nonimplantable bone growth stimulators are covered as a durable medical equipment benefit:

[A. Ultrasonic osteogenesis stimulator (e.g., the Sonic Accelerated Fracture Healing System (SAFHS)) to accelerate healing of fresh fractures, fusions, or delayed unions at either of the following high-risk sites:
   (I) Fresh fractures, fusions, or delayed unions of the shaft (diaphysis) of the tibia that are open or segmental; or
   (II) Fresh fractures, fusions, or delayed unions of the scaphoid (carpal navicular);

B. Ultrasonic osteogenesis stimulator for non-unions, failed arthrodesis, and congenital pseudarthrosis (pseudarthrosis) of the appendicular skeleton if there has been no progression of healing for three (3) or more months despite appropriate fracture care; or

C. Direct current electrical bone-growth stimulator is covered for the following indications:
   (I) Delayed unions of fractures or failed arthrodesis at high-risk sites (i.e., open or segmental tibial fractures, carpal navicular fractures);
   (II) Non-unions, failed fusions, and congenital pseudarthrosis where there is no evidence of progression of healing for three (3) or more months despite appropriate fracture care; or

[III] Members who are at high risk for spinal fusion failure when any of the following criteria is met:
   (a) A multiple-level fusion entailing three (3) or more vertebrae (e.g., L3 to L5, L4 to S1, etc.);
   (b) Grade II or worse spondylolisthesis; or
   (c) One (1) or more failed fusions;

7. Contraception and Sterilization. All Food and Drug Administration- (FDA-) approved contraceptive methods, sterilization procedures, and patient education and counseling for all women with reproductive capacity;

8. Cardiac rehabilitation. An electrocardiographically-monitored program of outpatient cardiac rehabilitation (Phase II) is covered for specific criteria when it is individually prescribed by a provider and a formal exercise stress test is completed following the event and prior to the initiation of the program. Cardiac rehabilitation is covered for members who meet one (1) of the following criteria:

[A. Acute myocardial infarction (MI) (heart attack in the last twelve (12) months);

B. Coronary artery bypass grafting (CABG);

C. Stable angina pectoris;

D. Percutaneous coronary vessel remodeling;

E. Valve replacement or repair;

F. Heart transplant;

G. Coronary artery disease (CAD) associated with chronic stable angina that has failed to respond adequately to pharmacotherapy and is interfering with the ability to perform age-related activities of daily living and/or impairing functional abilities; or

H. Heart failure that has failed to respond adequately to pharmacotherapy and is interfering with the ability to perform age-related activities of daily living and/or impairing functional abilities;

9. Chelation therapy. The administration of FDA-approved chelating agents is covered for any of the following conditions:

[A. Genetic or hereditary hemochromatosis;

B. Lead overload in cases of acute or long-term lead exposure;

C. Secondary hemochromatosis due to chronic iron overload due to transfusion-dependent anemias (e.g., Thalassemias, Cooley’s anemia, sickle cell anemia, sideroblastic anemia);

D. Copper overload in patients with Wilson’s disease;

E. Arsenic, mercury, iron, copper, or gold poisoning when long-term exposure to and toxicity has been confirmed through lab results or clinical findings consistent with metal toxicity;

F. Aluminum overload in chronic hemodialysis patients;

G. Emergency treatment of hypercalcemia;

H. Prophylaxis against doxorubicin-induced cardiomyopathy;
1. Internal plutonium, americium, or curium contamination; or
   J. Cystinuria;

10. Chiropractic services. Chiropractic—manipulation and adjunct therapeutic procedures/modalities [e.g., mobilization, therapeutic exercise, traction] are covered when all of the following conditions are met:
   [A. A neuromusculoskeletal condition is diagnosed that may be relieved by standard chiropractic treatment in order to restore optimal function;
   B. Chiropractic care is being performed by a licensed doctor of chiropractic who is practicing within the scope of his/her license as defined by state law;
   C. The individual is involved in a treatment program that clearly documents all of the following:
      (I) A prescribed treatment program that is expected to result in significant therapeutic improvement over a clearly defined period of time;
      (II) The symptoms being treated;
      (III) Diagnostic procedures and results;
      (IV) Frequency, duration, and results of planned treatment modalities;
      (V) Anticipated length of treatment plan with identification of quantifiable, attainable short-term and long-term goals; and
      (VI) Demonstrated progress toward significant functional gains and/or improved activity tolerances;
   D. Following previous successful treatment with chiropractic care, acute exacerbation or re-injury are covered when all of the following criteria are met:
      (I) The member reached maximal therapeutic benefit with prior chiropractic treatment;
      (II) The member was compliant with a self-directed homecare program;
      (III) Significant therapeutic improvement is expected with continued treatment; and
      (IV) The anticipated length of treatment is expected to be short-term [e.g., no more than six (6) visits within a three- (3-) week period];
   11. Clinical trials. Routine member care costs incurred as the result of a Phase I, II, III, or IV clinical trial that is conducted in relation to the prevention, detection, or treatment of cancer or other life-threatening disease or condition are covered when—
      A. The study or investigation is conducted under an investigational new drug application reviewed by the FDA; or
      B. Is a drug trial that is exempt from having such an investigational new drug application. Life-threatening condition means any disease or condition from which the likelihood of death is probable unless the course of the disease or condition is interrupted; and
      C. Routine member care costs include all items and services consistent with the coverage provided in plan benefits that would otherwise be covered for a member not enrolled in a clinical trial. Routine patient care costs do not include the investigational item, device, or service itself; items and services that are provided solely to satisfy data collection and analysis needs and are not used in the direct clinical management of the member; or a service that is clearly inconsistent with widely accepted and established standards of care for a particular diagnosis;
      D. The member must be eligible to participate in the clinical trial according to the trial protocol with respect to treatment of cancer or other life-threatening disease or condition; and
      E. The clinical trial must be approved or funded by one (1) of the following:
         (I) National Institutes of Health (NIH);
         (II) Centers for Disease Control and Prevention (CDC);
         (III) Agency for Health Care Research and Quality;
         (IV) Centers for Medicare & Medicaid Services (CMS);
         (V) A cooperative group or center of any of the previously named agencies or the Department of Defense or the Department of Veterans Affairs;
      (VI) A qualified non-governmental research entity identified in the guidelines issued by the National Institutes of Health for center support grants; or
      (VII) A study or investigation that is conducted by the Department of Veterans Affairs, the Department of Defense, or the Department of Energy and has been reviewed and approved to be comparable to the system of peer review of studies and investigations used by the NIH and assures unbiased review of the highest scientific standards by qualified individuals who have no interest in the outcome of the review;

12. Cochlear implant [device]. Uniaural (monaural) or binaural (bilateral) cochlear implantation and necessary replacement batteries are covered for a member with bilateral, pre- or post-linguistic, sensorineural, moderate-to-profound hearing impairment when there is reasonable expectation that a significant benefit will be achieved from the device and when the following age-specific criteria are met:
   and auditory brainstem implant;
   [A. Auditory brainstem implant. Auditory brainstem implant (ABI) covered for the diagnosis of neurofibromatosis type II, von Recklinghausen’s disease, or when a member is undergoing bilateral removal of tumors of the auditory nerves, and it is anticipated that the member will become completely deaf as a result of the surgery, or the member had bilateral auditory nerve tumors removed and is now bilaterally deaf;
      (I) For an adult (age eighteen (18) years or older) with BOTH of the following:
         (a) Bilateral, severe to profound sensorineural hearing loss determined by a pure-tone average of seventy (70) decibels (dB) hearing loss or greater at five hundred (500) hertz (Hz), one thousand (1000) Hz, and two thousand (2000) Hz; and
         (b) Member has limited benefit from appropriately fitted binaural hearing aids. Limited benefit from amplification is defined by test scores of forty percent (40%) correct or less in best-aided listening condition on open-set sentence cognition (e.g., Central Institute for the Deaf (CID) sentences, Hearing in Noise Test (HINT) sentences, and Consonant-Nucleus-Consonant (CNC) test);
      (II) For a child age twelve (12) months to seventeen (17) years, eleven (11) months with both of the following:
         (a) Profound, bilateral sensorineural hearing loss with thresholds of ninety (90) dB or greater at one thousand (1000) Hz; and
         (b) Limited or no benefit from a three- (3-) month trial of appropriately fitted binaural hearing aids;
      (III) For children four (4) years of age or younger, with one (1) of the following:
         (a) Failure to reach developmentally appropriate auditory milestones measured using the Infant-Toddler Meaningful Auditory Integration Scale, the Meaningful Auditory Integration Scale, or the Early Speech Perception test; or
         (b) Less than twenty percent (20%) correct on open-set word recognition test Multisyllabic Lexical Neighborhood Test (MLNLT) in conjunction with appropriate amplification and participation in intensive aural habilitation over a three- (3-) to six- (6-) month period;
      (IV) For children older than four (4) years of age with one (1) of the following:
         (a) Less than twelve percent (12%) correct on the Phonetically Balanced-Kindergarten Test; or
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...needed as a result of accident; and of disease. Treatment must be initiated within sixty (60) days of accident to sound natural teeth and tissue that are viable, functional, and free of disease...

B. Radiologic evidence of cochlear ossification;
C. The following additional medical necessity criteria must also be met for uniaural (monaural) or binaural (bilateral) cochlear implantation in adults and children:
   (I) Member must be enrolled in an educational program that supports listening and speaking with aided hearing;
   (II) Member must have had an assessment by an audiologist and from an otolaryngologist experienced in this procedure indicating the likelihood of success with this device;
   (III) Member must have no medical contraindications to cochlear implantation (e.g., cochlear aplasia, active middle ear infection); and
   (IV) Member must have arrangements for appropriate follow-up care, including the speech therapy required to take full advantage of this device;
D. A second cochlear implant is covered in the contralateral (opposite) ear as medically necessary in an individual with an existing unilateral cochlear implant when the hearing aid in the contralateral ear produces limited or no benefit;
E. The replacement of an existing cochlear implant is covered when either of the following criteria is met:
   (I) Currently used component is no longer functional and cannot be repaired; or
   (II) Currently used component renders the implant recipient unable to adequately and/or safely perform his/her age-appropriate activities of daily living; and
F. Post-cochlear or ABI rehabilitation program (aural rehabilitation) is covered to achieve benefit from a covered device;

A. Dental care is covered for the following:
   (I) Treatment to reduce trauma and restorative services limited to dental implants only when the result of accidental injury to sound natural teeth and tissue that are viable, functional, and free of disease. Treatment must be initiated within sixty (60) days of accident; and
   (II) Restorative services limited to dental implants when needed as a result of cancerous or non-cancerous tumors and cysts, cancer, and post-surgical sequelae.
B. The administration of general anesthesia, monitored anesthesia care, and hospital charges for dental care are covered for children younger than five (5) years, the severely disabled, or a person with a medical or behavioral condition that requires hospitalization when provided in a network or non-network hospital or surgical center;

14. Diabetes Self-Management Education;
15. Dialysis is covered when received through a network provider;
16. Durable medical equipment (DME) is covered when ordered by a provider to treat an injury or illness. DME includes, but is not limited to, the following:
   A. Insulin pumps;
   B. Oxygen;
   C. Augmentative communication devices;
   D. Manual and powered mobility devices;
E. Disposable supplies that do not withstand prolonged use and are periodically replaced, including, but not limited to, the following:
   (I) Colostomy and ureterostomy bags;
   (II) Prescription compression stockings limited to two (2) pairs or four (4) individual stockings per plan year;
   (III) Blood pressure cuffs/monitors with a diagnosis of diabetes;
   (IV) Repair and replacement of DME is covered when any of the following criteria are met:
      (I) Repairs, including the replacement of essential accessories, which are necessary to make the item or device serviceable;
      (II) Routine wear and tear of the equipment renders it non-functional and the member still requires the equipment; or
      (III) The provider has documented that the condition of the member changes or if growth-related;

17. Emergency room services. Coverage is for emergency medical conditions. If a member is admitted to the hospital, s/he may be required to transfer to network facility for maximum benefit. Hospital and ancillary charges are paid as a network benefit;
18. Eye glasses and contact lenses. Coverage limited to charges incurred in connection with the fitting of eye glasses or contact lenses for initial placement within one (1) year following cataract surgery;
19. Foot care (trimming of nails, corns, or calluses). Foot care services are covered when administered by a provider and—
   A. When associated with systemic conditions that are significant enough to result in severe circulatory insufficiency or areas of desensitization in the lower extremities including, but not limited to, any of the following:
      (I) Diabetes mellitus;
      (II) Peripheral vascular disease; or
      (III) Peripheral neuropathy.
   (IV) Evaluation/debridement of mycotic nails, in the absence of a systemic condition, when both of the following conditions are met:
      (a) Pain or secondary infection resulting from the thickening and dystrophy of the infected toenail plate; and
      (b) If the member is ambulatory, pain markedly limits ambulation;
20. Genetic counseling. Pre-test and post-test genetic counseling with a provider or a licensed or certified genetic counselor are covered when a member is recommended for covered heritable genetic testing. 
[A. Genetic counseling in connection with pregnancy management is covered only for evaluation of any of the following:
   (I) Couples who are closely related genetically (e.g., consanguinity, incest);
   (II) Familial cancer disorders;
   (III) Individuals recognized to be at increased risk for genetic disorders;
   (IV) Infertility cases where either parent is known to have a chromosomal abnormality;
   (V) Primary amenorrhea, azoopermia, abnormal sexual development, or failure in developing secondary sexual characteristics;
   (VI) Mother is a known, or presumed carrier of an X-linked recessive disorder;
   (VII) One (1) or both parents are known carriers of an autosomal recessive disorder;
   (VIII) Parents of a child born with a genetic disorder, birth defect, inborn error of metabolism, or chromosome abnormality;
   (IX) Parents of a child with intellectual developmental disorders, autism, developmental delays, or learning disabilities;
   (X) Pregnant women who, based on prenatal ultrasound tests or an abnormal multiple marker screening test,
maternal serum alpha-fetoprotein (AFP) test, test for sickle cell anemia, or tests for other genetic abnormalities have been told their pregnancy may be at increased risk for complications or birth defects;

[(XII) Pregnant women age thirty-five (35) years or older at delivery;

(XIII) Pregnant women, or women planning pregnancy, exposed to potentially teratogenic, mutagenic, or carcinogenic agents such as chemicals, drugs, infections, or radiation;

(XIV) When contemplating pregnancy, either parent affected with an autosomal dominant disorder;]

21. Genetic testing.

A. Genetic testing is covered to establish a molecular diagnosis of an inheritable disease when all of the following criteria are met:

(I) The member displays clinical features or is at direct risk of inheriting the mutation in question (pre-symptomatic);

(II) The result of the test will directly impact the treatment being delivered to the member;

(III) The testing method is considered scientifically valid for identification of a genetically-linked inheritable disease; and

(IV) After history, physical examination, pedigree analysis, genetic counseling, and completion of conventional diagnostic studies, a definitive diagnosis remains uncertain.

B. Genetic testing for the breast cancer susceptibility gene (BRCA) when family history is present;

22. Hair analysis. Chemical hair analysis is covered for the diagnosis of suspected chronic arsenic poisoning. Other purposes are considered experimental and investigational;

23. Hair prostheses. Prostheses and expenses for scalp hair prostheses worn for hair loss are covered for alopecia areata or alopecia totalis for children eighteen (18) years of age or younger. The annual maximum is two hundred dollars ($200), and the lifetime maximum is three thousand two hundred dollars ($3,200);

24. Hearing aids (per ear). Hearing aids [are] covered once every two (2) years for conductive hearing loss unresponsive to medical or surgical interventions, sensorineural hearing loss, and mixed hearing loss.

[A. Prior to receiving a hearing aid members must receive—

(I) A medical exam by a physician or other qualified provider to identify any medically treatable conditions that may affect hearing; and

(II) A comprehensive hearing test to assess the need for hearing aids conducted by a certified audiologist, hearing instrument specialist, or other provider licensed or certified to administer this test.

B. Covered once every two (2) years.] If the cost of one (1) hearing aid exceeds the amount listed below, member is also responsible for charges over that amount.

[(I)]

[A. Conventional: one thousand dollars ($1,000).

[B. Programmable: two thousand dollars ($2,000).

C. Digital: two thousand five hundred dollars ($2,500).

D. Bone Anchoring Hearing Aid (BAHA): three thousand five hundred dollars ($3,500);

25. Hearing testing. One (1) hearing test per year. Additional hearing tests are covered if recommended by provider;

26. Home health care. Skilled home health care is covered for members who are homebound because of injury or illness (i.e., the member leaves home only with considerable and taxing effort, and absences from home are infrequent or of short duration, or to receive medical care). Services must be performed by a registered nurse or licensed practical nurse, licensed therapist, or a registered dietitian. Covered services include:

A. Home visits instead of visits to the provider’s office that do not exceed the usual and customary charge to perform the same service in a provider’s office;

B. Intermittent nurse services. Benefits are paid for only one (1) nurse at any one (1) time, not to exceed four (4) hours per twenty-four (24)- hour period;

C. Nutrition counseling provided by or under the supervision of a registered dietitian;

D. Physical, occupational, respiratory, and speech therapy provided by or under the supervision of a licensed therapist;

E. Medical supplies, drugs, or medication prescribed by provider, and laboratory services to the extent that the plan would have covered them under this plan if the covered person had been in a hospital;

F. A home health care visit is defined as—

(I) A visit by a nurse providing intermittent nurse services (each visit includes up to a four- (4-) hour consecutive visit in a twenty-four (24-) hour period if clinical eligibility for coverage is met) or a single visit by a therapist or a registered dietitian; and

G. Benefits cannot be provided for any of the following:

(I) Homemaker or housekeeping services;

(II) Supportive environment materials such as handrails, ramps, air conditioners, and telephones;

(III) Services performed by family members or volunteer workers;

(IV) “Meals on Wheels” or similar food service;

(V) Separate charges for records, reports, or transportation;

(VI) Expenses for the normal necessities of living such as food, clothing, and household supplies; and

(VII) Legal and financial counseling services, unless otherwise covered under this plan;

27. Hospice care and palliative services (inpatient or outpatient). Includes bereavement and respite care. Hospice care services, including pre-hospice evaluation or consultation, are covered when the individual is terminally ill and expected to live six (6) months or less, potentially curative treatment for the terminal illness is not part of the prescribed plan of care, the individual or appointed designee has formally consented to hospice care (i.e., care directed mostly toward palliative care and symptom management), and the hospice services are provided by a certified/accredited hospice agency with care available twenty-four (24) hours per day, seven (7) days per week.]

[A. When the above criteria are met, the following hospice care services are covered:

(I) Assessment of the medical and social needs of the terminally ill person, and a description of the care to meet those needs;

(II) Inpatient care in a facility when needed for pain control and other acute and chronic symptom management, psychological and dietary counseling, physical or occupational therapy, and part-time home health care services;

(III) Outpatient care for other services as related to the terminal illness, which include services of a physician, physical or occupational therapy, and nutrition counseling provided by or under the supervision of a registered dietitian; and

(IV) Bereavement counseling benefits which are received by a member’s close relative when directly connected to the member’s death and bundled with other hospice charges. The services must be furnished within twelve (12) months of death.]

28. Hospital (includes inpatient, outpatient, and surgical centers).

A. The following benefits are covered:
(I) Semi-private room and board. For network charges, this rate is based on network repricing. For non-network charges, any charge over a semi-private room charge will be a covered expense only when clinical eligibility for coverage is met. If the hospital has no semi-private rooms, the plan will allow the private room rate subject to usual, customary, and reasonable charges or the network rate, whichever is applicable;

(II) Intensive care unit room and board;

(III) Surgery, therapies, and ancillary services including, but not limited to:

(a) Cornea transplant;

(b) Coverage for breast reconstruction surgery or prostheses following mastectomy and lumpectomy is available to both females and males. A diagnosis of breast cancer is not required for breast reconstruction services to be covered, and the timing of reconstructive services is not a factor in coverage;

(c) Sterilization for the purpose of birth control is covered;

(d) Cosmetic/reconstructive surgery is covered to repair a functional disorder caused by disease or injury;

(e) Cosmetic/reconstructive surgery is covered to repair a congenital defect or abnormality for a member younger than nineteen (19) years; and

(f) Blood, blood plasma, and plasma expanders are covered, when not available without charge;

(IV) Inpatient mental health services [are covered when authorized by a physician for treatment of a mental health disorder. Inpatient mental health services are covered, subject to all of the following:]; and

[(a) Member must be ill in more than one (1) area of daily living to such an extent that s/he is rendered dysfunctional and requires the intensity of an inpatient setting for treatment. Without such inpatient treatment, the member’s condition would deteriorate;

(b) The member’s mental health disorder must be treatable in an inpatient facility;

(c) The member’s mental health disorder must meet diagnostic criteria as described in the most recent edition of the American Psychiatric Association Diagnostic and Statistical Manual (DSM). If outside of the United States, the member’s mental health disorder must meet diagnostic criteria established and commonly recognized by the medical community in that region;

(d) The attending provider must be a psychiatrist. If the admitting provider is not a psychiatrist, a psychiatrist must be attending to the member within twenty-four (24) hours of admittance. Such psychiatrist must be United States board-eligible or board-certified. If outside of the United States, inpatient services must be provided by an individual who has received a diploma from a medical school recognized by the government agency in the country where the medical school is located. The attending provider must meet the requirements, if any, set out by the foreign government or regionally-recognized licensing body for treatment of mental health disorders;

(e) Day treatment (partial hospitalization) for mental health services means a day treatment program that offers intensive, multidisciplinary services provided on less than a full-time basis. The program is designed to treat patients with serious mental or nervous disorders and offers major diagnostic, psychosocial, and prevocational modalities. Such programs must be a less-restrictive alternative to inpatient treatment; and

(f) Mental health services received in a residential treatment facility that is licensed by the state in which it operates and provides treatment for mental health disorders is covered. This does not include services provided at a group home. If outside of the United States, the residential treatment facility must be licensed or approved by the foreign government or an accreditation or licensing body working in that foreign country; and]

(V) Outpatient mental health services [are covered if the member is at a therapeutic medical or mental health facility and treatment includes measurable goals and continued progress toward functional behavior and termination of treatment. Continued coverage may be denied when positive response to treatment is not evident. Treatment must be provided by one (1) of the following:];

[(a) A United States board-eligible or board-certified psychiatrist licensed in the state where the treatment is provided;

(b) A therapist with a doctorate or master’s degree that denotes a specialty in psychiatry [Psy.D.];

(c) A state-licensed psychologist;

(d) A state-licensed or certified social worker practicing within the scope of his or her license or certification; or

(e) Licensed professional counselor;]

29. Infusions are covered when received through a network provider. Medications (specialty and non-specialty) that can be safely obtained through a pharmacy and which may be self-administered are not a medical plan benefit but are covered as part of the pharmacy benefit;

30. Injections. See preventive services for coverage of vaccinations. See contraception and sterilization for coverage of birth control injections. Medications (specialty and non-specialty) that can be safely obtained through a pharmacy and which may be self-administered are not a medical plan benefit but are covered as part of the pharmacy benefit;]

[A. B12 injections are covered for the following conditions:

(I) Pernicious anemia;

(II) Crohn’s disease;

(III) Ulcerative colitis;

(IV) Inflammatory bowel disease;

(V) Intestinal malabsorption;

(VI) Fish tapeworm anemia;

(VII) Vitamin B12 deficiency;

(VIII) Other vitamin B12 deficiency anemia;

(IX) Macrocytic anemia;

(X) Other specified megaloblastic anemias;

(XI) Megaloblastic anemia;

(XII) Malnutrition of alcoholism;

(XIII) Thrombocytopenia, unspecified;

(XIV) Dementia in conditions classified elsewhere;

(XV) Polyneuropathy in diseases classified elsewhere;

(XVI) Alcoholic polyneuropathy;

(XVII) Regional enteritis of small intestine;

(XVIII) Postgastric surgery syndromes;

(XIX) Other prophylactic chemo-therapy;

(xx) Intestinal bypass or anastomosis status;

(XXI) Acquired absence of stomach;

(XXII) Pancreatic insufficiency; and

(XXIII) Ideopathic progressive polyneuropathy;]

31. Lab, X-ray, and other diagnostic procedures. Outpatient diagnostic services are covered when tests or procedures are performed for a specific symptom and to detect or monitor a condition. Professional charges for automated lab services performed by an out-of-network provider are not covered;

32. Maternity coverage. Prenatal and postnatal care is covered.
Route prenatal office visits and screenings recommended by the Health Resources and Services Administration are covered at one hundred percent (100%). Other care is subject to [the] applicable copayments, deductible, and coinsurance. Newborns and their mothers are allowed hospital stays of at least forty-eight (48) hours after vaginal birth and ninety-six (96) hours after cesarean section birth. If discharge occurs earlier than specific time periods, the plan shall provide coverage for post discharge care that shall consist of a two- (2-) visit minimum, at least one (1) in the home.

33. Nutritional counseling. Individualized nutritional evaluation and counseling for the management of any medical condition for which appropriate diet and eating habits are essential to the overall treatment program is covered when ordered by a physician or physician extender and provided by a licensed health-care professional (e.g., a registered dietitian);

34. Nutrition therapy.

(A) Nutrition therapy is covered only when the following criteria are met:

(I) Nutrition therapy is the sole source of nutrients or a significant percentage of the daily caloric intake;

(II) Nutrition therapy is used in the treatment of, or in association with, a demonstrable disease, condition, or disorder;

(III) Nutrition therapy is necessary to sustain life or health;

(IV) Nutrition therapy is prescribed by a provider; and

(V) Nutrition therapy is managed, monitored, and evaluated on an on-going basis, by a provider.

B. Only the following types of nutrition therapy are covered:

(I) Enteral Nutrition (EN). EN is the provision of nutritional requirements via the gastrointestinal tract. EN can be taken orally or through a tube into the stomach or small intestine;

(II) Parenteral Nutrition Therapy (PN) and Total Parenteral Nutrition (TPN). PN is liquid nutrition administered through a vein to provide part of daily nutritional requirements. TPN is a type of PN that provides all daily nutrient needs. PN or TPN are covered when the member’s nutritional status cannot be adequately maintained on oral or enteral feedings;

(III) Intradialytic Parenteral Nutrition (IDPN). IDPN is a type of PN that is administered to members on chronic hemodialysis during dialysis sessions to provide most nutrient needs. IDPN is covered when the member is on chronic hemodialysis and nutritional status cannot be adequately maintained on oral or enteral feedings;

35. Office visit. Member encounter with a provider for health care, mental health, or substance use disorder in an office, clinic, or ambulatory care facility is covered based on the service, procedure, or related treatment plan;

36. Oral surgery is covered for injury, tumors, or cysts. Oral surgery includes, but is not limited to, reduction of fractures and dislocation of the jaws; external incision and drainage of cellulites; incision of accessory sinuses, salivary glands, or ducts; excision of exostosis of jaws and hard palate; and frenectomy. Treatment must be initiated within sixty (60) days of accident. No coverage for dental care, including oral surgery, as a result of poor dental hygiene. Extractions of bony or partial bony impactions are excluded;

37. Orthognathic or Jaw Surgery. Orthognathic or jaw surgery is covered when one (1) of the following conditions is documented and diagnosed:

A. Acute traumatic injury, and post-surgical sequela;

B. [Cancerous or non-cancerous T]umors and cysts, cancer, and post-surgical sequela;

C. Cleft lip/palate (for cleft lip/palate related jaw surgery); or

D. Physical [or physiological] abnormality [when one (1) of the following criteria is met];

(ii) Anteroposterior Discrepancies—

(a) Maxillary/Mandibular incisor relationship: over jet of 5mm or more, or a 0 to a negative value (norm 2mm);

(b) Maxillary/Mandibular anteroposterior molar relationship discrepancy of 4mm or more (norm 0 to 1mm); or

(c) These values represent two (2) or more standard deviation from published norms;

(iii) Vertical Discrepancies—

(a) Presence of a vertical facial skeletal deformity which is two (2) or more standard deviations from published norms for accepted skeletal landmarks;

(b) Open bite with no vertical overlap of anterior teeth or unilateral or bilateral posterior open bite greater than 2mm;

(c) Deep overbite with impingement or irritation of buccal or lingual soft tissues of the opposing arch; or

(d) Supraeruption of a dentoalveolar segment due to lack of occlusion;

(iii) Transverse Discrepancies—

(a) Presence of a transverse skeletal discrepancy which is two (2) or more standard deviations from published norms; or

(b) Total bilateral maxillary palatal cusps to mandibular fossa discrepancy of 4mm or greater, or a unilateral discrepancy of 3mm or greater, given normal axial inclination of the posterior teeth; or

(iv) Asymmetries—

(a) Anteroposterior, transverse, or lateral asymmetries greater than 3mm with concomitant occlusal asymmetry;

(v) Masticatory (chewing) and swallowing dysfunction due to malocclusion (e.g., inability to incise or chew solid foods, choking on incompletely masticated solid foods, damage to soft tissue during mastication, malnutrition);

(vi) Speech impairment; or

(vii) Obstructive sleep apnea or airway dysfunction;

38. Orthotics.

A. Ankle–Foot Orthosis (AFO) and Knee–Ankle–Foot Orthosis (KAFO).

(I) Basic coverage criteria for AFO and KAFO used during ambulation are as follows:

(a) AFO is covered when used in ambulation for members with weakness or deformity of the foot and ankle, which require stabilization for medical reasons, and have the potential to benefit functionally;

(b) KAFO is covered when used in ambulation for members when the following criteria are met:

I. Member is covered for AFO; and

II. Additional knee stability is required; and

(c) AFO and KAFO that are molded-to-patient-model, or custom-fabricated, are covered when used in ambulation, only when the basic coverage criteria and one (1) of the following criteria are met:

I. The member could not be fitted with a prefabricated AFO;

II. AFO or KAFO is expected to be permanent or for more than six (6) months duration;

III. Knee, ankle, or foot must be controlled in more than one (1) plane;

IV. There is documented neurological, circulatory, or orthopedic status that requires custom fabricating over a model to prevent tissue injury; or

V. The member has a healing fracture which lacks normal anatomical integrity or anthropometric proportions.

AFO and KAFO Not Used During Ambulation.
(a) AFO and KAFO not used in ambulation are covered if the following criteria are met:
   I. Passive range of motion test was measured with agoniometer and documented in the medical record;
   II. Documentation of an appropriate stretching program administered under the care of provider or caregiver;
   III. Plantar flexion contracture of the ankle with dorsiflexion on passive range of motion testing of at least ten degrees (10°) (i.e., a non-fixed contracture);
   IV. Reasonable expectation of the ability to correct the contracture;
   V. Contracture is interfering or expected to interfere significantly with the patient’s functional abilities; and
   VI. Used as a component of a therapy program which includes active stretching of the involved muscles and/or tendons; or
   VII. Member has plantar fasciitis.

(b) Replacement interface for AFO or KAFO is covered only if member continues to meet coverage criteria and is limited to a maximum of one (1) per six (6) months.

B. Cast Boot, Post-Operative Sandal or Shoe, or Healing Shoe. A cast boot, post-operative sandal or shoe, or healing shoe is covered for one (1) of the following indications:
   (I) To protect a cast from damage during weight-bearing activities following injury or surgery;
   (II) To provide appropriate support and/or weight-bearing surface to a foot following surgery;
   (III) To promote good wound care and/or healing via appropriate weight distribution and foot protection; or
   (IV) When the patient is currently receiving treatment for lymphedema and the foot cannot be fitted into conventional footwear.

C. Cranial Orthoses. Cranial orthosis is covered for Synostotic and Non-Synostotic Plagiocephaly. Plagiocephaly is an asymmetrically shaped head. Synostotic Plagiocephaly is due to premature closure of cranial sutures. Non-Synostotic Plagiocephaly is from positioning or deformation of the head. Cranial orthosis is the use of a special helmet or band on the head which aids in molding the shape of the cranium to normal. Initial reimbursement shall cover any subsequent revisions.

D. Elastic Supports. Elastic supports are covered when prescribed for one (1) of the following indications:
   (I) Severe or incapacitating vascular problems, such as acute thrombophlebitis, massive venous stasis, or pulmonary embolism;
   (II) Venous insufficiency;
   (III) Varicose veins;
   (IV) Edema of lower extremities;
   (V) Edema during pregnancy; or
   (VI) Lymphedema.

E. Footwear Incorporated Into a Brace for Members with Skeletally Mature Feet. Footwear incorporated into a brace must be billed by the same supplier billing for the brace. The following types of footwear incorporated into a brace are covered:
   (I) Orthopedic footwear;
   (II) Other footwear such as high top, depth inlay, or custom;
   (III) Heel replacements, sole replacements, and shoe transfers involving shoes on a brace;
   (IV) Inserts for a shoe that is an integral part of a brace and are required for the proper functioning of the brace; or
   (V) Other shoe modifications if they are on a shoe that is an integral part of a brace and are required for the proper functioning of the brace.

F. Foot Orthoses. Custom, removable foot orthoses are covered [for members who meet the following criteria]:
   (I) Member with skeletally mature feet who has any of the following conditions:
      (a) Acute plantar fasciitis;
      (b) Acute sport-related injuries with diagnoses related to inflammatory problems such as bursitis or tendinitis;
      (c) Calcaneal bursitis (acute or chronic);
      (d) Calcaneal spurs (HEEL spurs);
      (e) Conditions related to diabetes;
      (f) Inflammatory conditions (e.g., sepsis, submetatarsal bursitis, synovitis, tenosynovitis, sylvial cyst, osteomyelitis, and plantar fascial fibromatosis);
      (g) Medial osteoarthritis of the knee;
      (h) Musculoskeletal/arthropathic deformities including deformities of the joint or skeleton that impairs walking in a normal shoe (e.g., bunions, hallux valgus, talipes deformities, pes deformities, or anomalies of toes);
      (i) Neurologically impaired feet including neuro-ma, tarsal tunnel syndrome, ganglionic cyst;
      (j) Neuropathies involving the feet, including those associated with peripheral vascular disease, diabetes, carcinoma, drugs, toxins, and chronic renal disease; or
      (k) Vascular conditions including ulceration, poor circulation, peripheral vascular disease, Buerger’s disease (thromboangiitis obliterans), and chronic thrombophlebitis;
   (II) Member with skeletally immature feet who has any of the following conditions:
      (a) Hallux valgus deformities;
      (b) In-toe or out-toe gait;
      (c) Musculoskeletal weakness such as pronation or pes planus;
      (d) Structural deformities such as tarsal coalition;
      (e) Torsional conditions such as metatarsus adductus, tibial torsion, or femoral torsion.

G. Helmets. Helmets are covered when cranial protection is required due to a documented medical condition that makes the member susceptible to injury during activities of daily living.

H. Hip Orthosis. Hip orthosis is covered for one (1) of the following indications:
   (I) To reduce pain by restricting mobility of the hip;
   (II) To facilitate healing following an injury to the hip or related soft tissues;
   (III) To facilitate healing following a surgical procedure of the hip or related soft tissue; or
   (IV) To otherwise support weak hip muscles or a hip deformity.

I. Knee Orthosis. Knee orthosis is covered for one (1) of the following indications:
   (I) To reduce pain by restricting mobility of the knee;
   (II) To facilitate healing following an injury to the knee or related soft tissues;
   (III) To facilitate healing following a surgical procedure on the knee or related soft tissue; or
   (IV) To otherwise support weak knee muscles or a knee deformity.

J. Orthopedic Footwear for Diabetic Members.

   (I) Orthopedic footwear, therapeutic shoes, inserts, or modifications to therapeutic shoes are covered for diabetic members if any following criteria are met:
      (a) Previous amputation of the other foot or part of either foot;
      (b) History of previous foot ulceration of either foot;
      (c) History of pre-ulcerative calluses of either foot;
      (d) Peripheral neuropathy with evidence of callus formation of either foot;
      (e) Foot deformity of either foot; or
      (f) Poor circulation in either foot.

   (II) Coverage is limited to one (1) of the following within one (1) year:
      (a) One (1) pair of custom molded shoes (which includes
inserts provided with these shoes) and two (2) additional pairs of inserts;

(b) One (1) pair of depth shoes and three (3) pairs of inserts (not including the non-customized removable inserts provided with such shoes); or

(c) Up to three (3) pairs of inserts not dispensed with diabetic shoes if the supplier of the shoes verifies in writing that the patient has appropriate footwear into which the insert can be placed.

K. Orthotic-Related Supplies. Orthotic-related supplies are covered when necessary for the function of the covered orthotic device.

L. Spinal Orthoses. A thoracic-lumbar-sacral orthosis, lumbar orthosis, lumbar-sacral orthosis, and cervical orthosis are covered for the following indications:

(I) To reduce pain by restricting mobility of the trunk;
(II) To facilitate healing following an injury to the spine or related soft tissues;
(III) To facilitate healing following a surgical procedure of the spine or related soft tissue; or
(IV) To otherwise support weak spinal muscles or a deformed spine.

M. Trusses. Trusses are covered when a hernia is reducible with the application of a truss.

N. Upper Limb Orthosis. Upper limb orthosis is covered for the following indications:

(I) To reduce pain by restricting mobility of the joint(s); or
(II) To facilitate healing following an injury to the joint(s) or related soft tissues; or
(III) To facilitate healing following a surgical procedure of the joint(s) or related soft tissue.

O. Orthotic Device Replacement. When repairing an item that is no longer cost-effective and is out of warranty, the plan will consider replacing the item subject to review of medical necessity and life expectancy of the device.

39. Preventive services.
A. Services recommended by the U.S. Preventive Services Task Force (categories A and B).
B. Vaccinations recommended by the Advisory Committee on Immunization Practices of the Centers for Disease Control and Prevention.
C. Preventive care and screenings for infants, children, and adolescents supported by the Health Resources and Services Administration.
D. Preventive care and screenings for women supported by the Health Resources and Services Administration.
E. Preventive exams and other services ordered as part of the exam. For benefits to be covered as preventive, they must be coded by the provider as routine, without indication of an injury or illness.
F. Cancer screenings. One (1) per calendar year. Additional screenings beyond one (1) per calendar year covered as diagnostic unless otherwise specified—

(I) Mammograms—no age limit. Standard two-dimensional (2D) breast mammography and breast tomosynthesis (three-dimensional (3D) mammography);
(II) Pap smears—no age limit;
(III) Prostate—no age limit; and
(IV) Colorectal screening—no age limit.

G. Online weight management program offered through the plan’s exclusive provider arrangement;

40. Prostheses (prosthetic devices). Basic equipment that meets medical needs. Repair and replacement is covered due to normal wear and tear, if there is a change in medical condition, or if growth-related;

41. Pulmonary rehabilitation. Comprehensive, individualized, goal-directed outpatient pulmonary rehabilitation covered for pre-and post-operative intervention for lung transplantation and lung volume reduction surgery (LVRS) or when all of the following apply:

A. Member has a reduction of exercise tolerance that restricts the ability to perform activities of daily living (ADL) or work;
B. Member has chronic pulmonary disease (including asthma, emphysema, chronic bronchitis, chronic airflow obstruction, cystic fibrosis, alpha-1 antitrypsin deficiency, pneumonia, tuberculosis, radiation pneumonitis, pulmonary fibrosis, pulmonary alveolar proteinosis, pulmonary hemosiderosis, fibrosing alveolitis), or other conditions that affect pulmonary function such as ankylosing spondylitis, scoliosis, myasthenia gravis, muscular dystrophy, Guillain-Barré syndrome, or other infective polyneuritis, sarcoidosis, paralysis of diaphragm, or bronchopulmonary dysplasia; and

C. Member has a moderate to moderately severe functional pulmonary disability, as evidenced by either of the following, and does not have any concomitant medical condition that would otherwise iniminate contribute to deterioration of pulmonary status or undermine the expected benefits of the program (e.g., symptomatic coronary artery disease, congestive heart failure, myocardial infarction within the last six (6) months, dysrhythmia, active joint disease, claudication, malignancy):

(I) A maximal pulmonary exercise stress test under optimal bronchodilatory treatment which demonstrates a respiratory limitation to exercise with a maximal oxygen uptake (VO2,max) equal to or less than twenty milliliters per kilogram per minute (20 mL/kg/min), or about five (5) metabolic equivalents (METS); or
(II) Pulmonary function tests showing that either the Forced Expiratory Volume in One Second (FEV1), Forced Vital Capacity (FVC), FEV1/FVC, or Diffusing Capacity of the Lung for Carbon Monoxide (DLCO) is less than sixty percent (60%) of that predicted;

42. Skilled Nursing Facility. Skilled nursing facility services are covered up to one hundred twenty (120) days per calendar year;

43. Telehealth Services. Telehealth services are covered for the diagnosis, consultation, or treatment of a member on the same basis that the service would be covered when it is delivered in person;

44. Therapy. Physical, occupational, and speech therapy are covered when prescribed by a provider and subject to the provisions below:

A. Physical therapy.

(I) Physical therapy must meet the following criteria:

(a) The program is designed to improve lost or impaired physical function or reduce pain resulting from illness, injury, congenital defect, or surgery;
(b) The program is expected to result in significant therapeutic improvement over a clearly defined period of time; and
(c) The program is individualized, and there is documentation outlining quantifiable, attainable treatment goals;

B. Occupational therapy must meet the following criteria:

(I) The program is designed to improve or compensate for lost or impaired physical functions, particularly those affecting activities of daily living, resulting from illness, injury, congenital defect, or surgery;

(II) The program is expected to result in significant therapeutic improvement over a clearly defined period of time; and
(III) The program is individualized, and there is documentation outlining quantifiable, attainable treatment goals;

C. Speech therapy.

(I) All of the following criteria must be met for coverage of speech therapy:

(a) The therapy requires one-to-one intervention and supervision of a speech-language pathologist;
(b) The therapy plan includes specific tests and measures that will be used to document significant progress every two (2) weeks;
(c) Meaningful improvement is expected;
(d) The therapy includes a transition from one-to-one supervision to a self- or caregiver- provided maintenance program upon discharge; and
(e) One (1) of the following:
I. Member has severe impairment of speech-language; and an evaluation has been completed by a certified speech-language pathologist that includes age-appropriate standardized tests to measure the extent of the impairment, performance deviation, and language and pragmatic skill assessment levels; or

II. Member has a significant voice disorder that is the result of anatomic abnormality, neurological condition, or injury (e.g., vocal nodules or polyps, vocal cord paresis or paralysis, post-operative vocal cord surgery);

45. Transplants. Stem cell, kidney, liver, heart, lung, pancreas, small bowel, or any combination are covered. Includes services related to organ procurement and donor expenses if not covered under another plan. Member must contact medical plan for arrangements.

A. Network includes travel and lodging allowance for the transplant recipient and an immediate family travel companion when the transplant facility is more than fifty (50) miles from the recipient’s residence. If the recipient is younger than age nineteen (19) years, travel and lodging is covered for both parents. The transplant recipient must be with the travel companion or parent(s) for the travel companion’s or parent(s’) travel expense to be reimbursable. Combined travel and lodging expenses are limited to a ten thousand dollar ($10,000) maximum per transplant.

(I) Lodging—maximum lodging expenses shall not exceed the per diem rates as established annually by U.S. General Services Administration (GSA) for a specific city or county. Go to www.gsa.gov for per diem rates.

(II) Travel—IRS standard medical mileage rates (same as flexible spending account (FSA) reimbursement).

(III) Meals—not covered.

B. Non-network. Charges above the maximum for services rendered at a non-network facility are the member’s responsibility and do not apply to the member’s deductible or out-of-pocket maximum. Travel, lodging, and meals are not covered;

46. Urgent care. Member encounter with a provider for urgent care is covered based on the service, procedure, or related treatment plan; and

47. Vision. One (1) routine exam and refraction is covered per calendar year.


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 2—State Membership

EMERGENCY AMENDMENT

22 CSR 10-2.061 Plan Limitations. The Missouri Consolidated Health Care Plan is amending section (1).

PURPOSE: This amendment revises the following benefits not payable by the plan: alternative therapies, assistant surgeon services, birthing center, home births, nocturnal enuresis alarm, therapy, vacuum extractions requested by a third party, and renumbers as necessary.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (employees, retirees, officers, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to members as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(1) Benefits shall not be payable for, or in connection with, any medical benefits, services, or supplies which do not come within the definition of covered charges. In addition, the items specified in this rule are not covered unless expressly stated otherwise and then only to the extent expressly provided herein or in 22 CSR 10-2.055 or 22 CSR 10-2.090.

(C) Alternative therapies—that are outside conventional medicine [including, but not limited to, acupuncture, acupressure, homeopathy, hypnosis, massage therapy, reflexology, and biofeedback] as determined by the claims administrator.

(E) Assistant surgeon services—unless determined to meet the clinical eligibility for coverage under the plan.

(F) Athletic enhancement services and sports performance training.

(G) Autopsy.

(H) Birthing center.

(I) Blood donor expenses.

(J) Blood pressure cuffs/monitors.

(K) Care received without charge.

(L) Charges exceeding the vendor contracted rate or benefit limit.

(M) Charges resulting from the failure to appropriately cancel a scheduled appointment.

(N) Childbirth classes.

(O) Comfort and convenience items.

(P) Cosmetic procedures.

(Q) Custodial or domiciliary care—including services and supplies that assist members in the activities of daily living such as walking, getting in and out of bed, bathing, dressing, feeding, and using the toilet; preparation of special diets; supervision of medication that is usually self-administered; or other services that can be performed by persons who are not providers.

(R) Dental care, including oral surgery.

(S) Devices or supplies bundled as part of a service are not
separately covered.

- **(TT)** Dialysis received through a non-network provider.
- **(UU)** Educational or psychological testing unless part of a treatment program for covered services.
- **(VV)** Examinations requested by a third party.
- **(WW)** Exercise equipment.
- **(XX)** Experimental/investigational/unproven services, procedures, supplies, or drugs as determined by the claims administrator.
- **(YY)** Eye exercises and associated expenses for orthoptics, eye exercises, radial keratotomy, LASIK, and other refractive eye surgery.
- **(ZZ)** Genetic testing based on family history alone, except for breast cancer susceptibility gene (BRCA) testing.
- **(AAA)** Health and athletic club membership—including costs of enrollment.
- **(BBB)** Hearing aid replacement batteries.
- **(CC)** Home births.
- **(DD)** Infertility treatment beyond the covered services to diagnose the condition.
- **(EE)** Infusions received through a non-network provider.
- **(FF)** Level of care, greater than is needed for the treatment of the illness or injury.
- **(GG)** Long-term care.
- **(HH)** Maxillofacial surgery.
- **(II)** Medical care and supplies to the extent that they are payable under—
  1. A plan or program operated by a national government or one of its agencies; or
  2. Any state’s cash sickness or similar law, including any group insurance policy approved under such law.
- **(JJ)** Medical service performed by a family member—including a person who ordinarily resides in the subscriber’s household or is related to the member, such as a spouse, parent, child, sibling, or brother/sister-in-law.
- **(KK)** Military service-connected injury or illness—including expenses relating to Veterans Affairs or a military hospital.
- **(LL)** Never events—never events on a list compiled by the National Quality Forum of inexcusable outcomes in a health care setting.
- **(MM)** Nocturnal enuresis alarm.
- **(NN)** Drugs that the pharmacy benefit manager (PBM) has excluded from the formulary and will not cover as a non-formulary drug unless it is approved in advance by the PBM.
- **(OO)** Non-medically necessary services.
- **(PP)** Non-provider allergy services or associated expenses relating to an allergic condition, including installation of air filters, air purifiers, or air ventilation system cleaning.
- **(QQ)** Non-reusable disposable supplies.
- **(RR)** Online weight management programs.
- **(SS)** Other charges as follows:
  1. Charges that would not otherwise be incurred if the subscriber was not covered by the plan; and
  2. Charges for which the subscriber or his/her dependents are not legally obligated to pay including, but not limited to, any portion of any charges that are discounted; and
  3. Charges made in the subscriber’s name but which are actually due to the injury or illness of a different person not covered by the plan; and
  4. No coverage for miscellaneous service charges including, but not limited to, charges for telephone consultations, administrative fees such as filling out paperwork or copy charges, or late payments.
- **(TT)** Over-the-counter medications with or without a prescription including, but not limited to, analgesics, antipyretics, non-sedating antihistamines, unless otherwise covered as a preventive service.
- **(UU)** Physical and recreational fitness.
- **(VV)** Private-duty nursing.
enhance job, school, or recreational performance; and
1. Long-term rehabilitative services when significant therapeutic improvement is not expected.

[CC] (XX) Travel expense and [ODD] Vaccinations requested by third party.

[EE] (YY) Workers’ Compensation services or supplies for an illness or injury eligible for, or covered by, any federal, state, or local government Workers’ Compensation Act, occupational disease law, or other similar legislation.


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 2—State Membership

EMERGENCY AMENDMENT

22 CSR 10-2.070 Coordination of Benefits. The Missouri Consolidated Health Care Plan is amending section (3).

PURPOSE: This amendment revises the order of benefit determination rules and renumbers as necessary.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (employees, retirees, officers, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to members as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(3) Order of Benefit Determination Rules.

(B) Rules. MCHCP determines its order of benefits [using the first of the following rules which applies] as follows:

1. Active/inactive employee. The benefits of the plan which covers the person as an employee who is neither laid off nor retired (or as that employee’s dependent) are determined before those of the plan which covers that person as a laid off or retired employee (or as that employee’s dependent);

2. Nondependent/dependent. The benefits of the plan which covers the person as an employer or subscriber (that is, other than as a dependent) are determined before those of the plan which covers the person as a dependent;

1. Non-Dependent/Dependent:

A. The plan which covers the member as an employee or subscriber is primary.

B. The plan which covers the member as dependent is secondary.

2. Active/layoff. The plan that covers the member or dependent through the member’s active employment is primary to a plan that covers the member or dependent through the member’s status as a laid off employee.

3. Retiree. The plan that covers the member or dependent through the member’s active employment is primary to a plan that covers the member or dependent through the member’s status as a retiree.


A. If a member is an active employee and has Medicare, MCHCP is the primary plan for the active employee and his/her dependents. Medicare is the secondary plan except for members with end stage renal disease (ESRD) as defined in subparagraph (3)(B)/3/4.D.

B. If a member is a retiree and has Medicare, Medicare is the primary plan for the retiree and his/her Medicare-eligible dependents. MCHCP is the secondary plan.

C. If a terminated vested employee with Medicare maintains coverage through one (1) of the MCHCP plans, Medicare is the primary plan and MCHCP is secondary.

D. If a member or his/her dependents are eligible for Medicare solely because of ESRD, the member’s MCHCP plan is primary to Medicare during the first thirty (30) months of Medicare eligibility for home peritoneal dialysis or home hemodialysis and thirty-three (33) months for in-center dialysis. After the thirty (30) or thirty-three (33) months, Medicare becomes primary, and claims are submitted first to Medicare, then to MCHCP for secondary coverage. The member is responsible for notifying MCHCP of his/her Medicare status.

E. If a member is on long-term disability through the Missouri State Employees’ Retirement System or the Public School Retirement System and is eligible for Medicare, Medicare is the primary plan and MCHCP plan is secondary;

/4/5. Dependent child/parents not separated or divorced. When MCHCP and another plan cover the same child as a dependent of different [persons, called] parents—

A. The benefits of the plan of the parent whose birthday falls earlier in a year are determined before those of the plan of the parent whose birthday falls later in that year; but

B. If both parents have the same birthday, the benefits of the plan which covered one (1) parent longer are determined before those of the plans which covered the other parent for a shorter period of time;

/5/6. Dependent child/separated, divorced, or never married. If two (2) or more plans cover a person as a dependent child of divorced, separated, or never married parents, benefits for the child are determined in this order—

A. First, the plan of the parent with custody of the child;

B. Then, the plan of the spouse of the parent with the custody
of the child;

C. Then, the plan of the parent not having custody of the child; and

D. Finally, the plan of the spouse of the parent not having custody of the child. However, if the specific terms of a court decree state that one (1) of the parents is responsible for the health care expense of the child and the entity obligated to pay or provide the benefits of the plan of that parent or spouse of the other parent has actual knowledge of those terms, the benefits of that plan are determined first. The plan of the other parent shall be the secondary plan. This paragraph does not apply with respect to any claim determination period or plan year during which any benefits are actually paid or provided before the entity has that actual knowledge;

[6./7.] Joint custody. If the specific terms of a court decree state that the parents shall share joint custody, without stating that one (1) of the parents is responsible for the health care expenses of the child, the plans covering the child shall follow the order of benefit determination rules outlined in paragraph (3)(B)/4./5.

[7./8.] Dependent child/parents both parents covered by MCHCP. If both parents are covered by MCHCP and both parents cover the child as a dependent, MCHCP will not coordinate benefits with itself;

[8./9.] (The plan that covers the member as a spouse is primary over the plan that covers the member as a dependent child) When an adult dependent is covered by both spouse and parent, the benefits of the plan which covered a person longer are determined before those of the plan which covered that person for the shorter term; and

[9./10.] Longer/shorter length of coverage. If none of the previous rules determines the order of benefits, the benefits of the plan which covered a person longer are determined before those of the plan which covered that person for the shorter term.


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Chapter 10—Health Care Plan
Chapter 2—State Membership

EMERGENCY AMENDMENT

22 CSR 10-2.075 Review and Appeals Procedure. The Missouri Consolidated Health Care Plan is amending sections (1), (3), and (5).

PURPOSE: This amendment revises the claim submission and initial benefit determinations time frames, updates the name and appeal contact information for the third party administrator.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (employees, retirees, officers, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or the availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to members as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(1) Claims Submissions and Initial Benefit Determinations PPO 750 Plan, PPO 1250 Plan, and Health Savings Account (HSA) Plan members.

(B) Medical and pharmacy service claims are divided into three (3) types: pre-service, post-service, and concurrent claims.

1. Pre-service claims are requests for approval that the plan or vendor requires a member to obtain before getting medical care or filling a prescription, such as prior authorization or a decision whether a treatment, procedure, or medication is medically necessary.

A. Pre-service claims must be decided within a reasonable period of time appropriate to the medical circumstances, but no later than [fifteen (15) days] twenty (20) business days from the date the vendor receives the claim. The vendor may extend the time period up to an additional [fifteen (15)] twenty (20) days if, for reasons beyond the vendor’s control, the decision cannot be made within the first [fifteen (15)] twenty (20) days. The vendor must notify the member prior to the expiration of the first fifteen- (15-) day period, explain the reason for the delay, and request any additional information. If more information is requested, the member has at least forty-five (45) days to provide the information to the vendor. The vendor then decide the claim no later than fifteen (15) days after the additional information is supplied or after the period of time allowed to supply it ends, whichever is first.

B. Urgent care claims are a special type of pre-service claim that require a quicker decision because waiting the standard time could seriously jeopardize the member’s life, health, or ability to regain maximum function. A request for an urgent care claim may be submitted verbally or in writing and will be decided within seventy-two (72) hours. Written confirmation of the decision will be sent by the vendor [as soon as possible thereafter] within three (3) business days.

2. Post-service claims are all other claims for services including claims after medical or pharmacy services have been provided, such as requests for reimbursement or payment of the costs for the services provided.

A. Post-service claims must be decided within a reasonable period of time, but not later than [thirty (30) days] (20) business days after the vendor receives the claim. If, because of reasons
beyond the vendor’s control, more time is needed to review the
claim, the vendor may extend the time period up to an additional [fifteen (15)/ thirty (30) days]. The vendor must notify the member
prior to the expiration of the first [fifteen- (15-) day/ twenty- (20-)
] day period, explain the reason for the delay, and request any addi-
tional information. If more information is requested, the member has at
least forty-five (45) days to provide the information to the vendor.
The vendor then must decide the claim no later than [fifteen (15)
days/ thirty (30) days] after the additional information is supplied or
after the period of time allowed to supply it ends, whichever is first.

3. Concurrent claims are claims related to an ongoing course of
previously-approved treatment. If the plan or vendor has approved
an ongoing course of treatment to be provided over a period of time or
number of treatments, any reduction or termination of the course of
treatment will be treated as a benefit denial. The plan or vendor will
notify a member in writing prior to reducing or ending a previously-
approved course of treatment in sufficient time to allow the member
or the member’s provider to appeal and obtain a determination before
the benefit is reduced or terminated.

(A) Definitions. Notwithstanding any other rule in this chapter to
the contrary, for purposes of a member’s right to appeal any adverse
benefit determination made by the plan, the plan administrator, a
claims administrator, or a medical or pharmacy benefit vendor, relat-
ing to the provision of health care benefits, other than those provided
in connection with the plan’s dental or vision benefit offering, the fol-
lowing definitions apply:

1. Adverse benefit determination. An adverse benefit determi-
nation means any of the following:
   A. A denial, reduction, or termination of, or a failure to
      provide or make payment (in whole or in part) for a benefit, includ-
      ing any denial, reduction, termination, or failure to provide or make
      payment that is based on a determination of an individual’s eligibility
      to participate in the plan;
   B. A denial, reduction, or termination of, or a failure to pro-
      vide or make payment (in whole or in part) for a benefit resulting
      from the application of any utilization review, as well as a failure to
      cover an item or service for which benefits are otherwise provided
      because it is determined to be experimental or investigational or not
      medically necessary or appropriate; or
   C. Any rescission of coverage after an individual has been
      covered under the plan;

2. Appeal (or internal appeal). An appeal or internal appeal
means review by the plan, the plan administrator, a claims adminis-
trator, or a medical or pharmacy benefit vendor of an adverse benefit
determination;

3. Claimant. Claimant means an individual who makes a claim
under this subsection. For purposes of this subsection, references to
claimant include a claimant’s authorized representative;

4. External review. The United States Department of Health and
Human Services (HHS) conducts external reviews for adverse benefit
determinations regarding medical and pharmacy benefits adminis-
tered by [UMR, Aetna Anthem, and Express Scripts, Inc. that
involve medical judgment (including, but not limited to), those based
on medical necessity, appropriateness, health care setting, level of
care, or effectiveness of a covered benefit; or a determination that a
treatment is experimental or investigational] and a rescission of cov-
erage (regardless of whether or not the rescission has any effect on
any particular benefit at that time);

5. Final internal adverse benefit determination. A final internal
adverse benefit determination means an adverse benefit determina-
tion that has been upheld by the plan, the plan administrator, a claims
administrator, or a medical or pharmacy benefit vendor at the com-
pletion of the internal appeals process under this subsection, or an
adverse benefit determination with respect to which the internal
appeals process has been deemed exhausted by application of applic-
able state or federal law;

6. Final external review decision. A final external review deci-
sion means a determination rendered under the external review
process at the conclusion of an external review; and

7. Recision. A rescission means a termination or discontinu-
amce of medical or pharmacy coverage that has retroactive effect
except that a termination or discontinuance of coverage is not a rescission if—
   A. The termination or discontinuance of coverage has only a
      prospective effect; or
   B. The termination or discontinuance of coverage is effective
      retroactively to the extent it is attributable to a failure to timely pay
      required premiums or contributions towards the cost of coverage.

(B) Internal Appeals.

1. Eligibility, termination for failure to pay, or rescission. Adverse
benefit determinations denying or terminating an individ-
ual’s coverage under the plan based on a determination of the indi-
vidual’s eligibility to participate in the plan or the failure to pay pre-
miums, or any rescission of coverage based on fraud or intentional
misrepresentation of a member or authorized representative of a
member are appealable exclusively to the Missouri Consolidated
Health Care Plan (MCHCP) Board of Trustees (board).

   A. The internal review process for appeals relating to eli-
      gibility, termination for failure to pay, or rescission shall consist of one
      (1) level of review by the board.
   B. Adverse benefit determination appeals to the board must
      identify the eligibility, termination, or rescission decision being
      appealed and the reason the claimant believes the MCHCP staff deci-
      sion should be overturned. The member should include with his/her
      appeal any information or documentation to support his/her appeal
      request.
   C. The appeal will be reviewed by the board in a meeting
closed pursuant to section 610.021, RSMo, and the appeal will be
      responded to in writing to the claimant within sixty (60) days from
      the date the board received the written appeal.

   D. Determinations made by the board constitute final internal
      adverse benefit determinations and are not eligible for external
      review except as specifically provided in 22 CSR 10-2.075(4)(A)4.

2. Medical and pharmacy services. Members may request inter-
 nal review of any adverse benefit determination relating to urgent
care, pre-service claims, and post-service claims made by the plan’s
medical and pharmacy vendors.

   A. Appeals of adverse benefit determinations shall be submit-
ted in writing to the vendor that issued the original determination
giving rise to the appeal at the applicable address set forth in this
rule.
   B. The internal review process for adverse benefit determina-
tions relating to medical services consists of two (2) levels of internal
review provided by the medical vendor that issued the adverse benefit
determination.

   (I) First level appeals must identify the decision being
   appealed and the reason the member believes the original claim deci-
   sion should be overturned. The member should include with his/her
   appeal any additional information or documentation to support the
   reason the original claim decision should be overturned.

   (II) First level appeals will be reviewed by the vendor by
   someone who was not involved in the original decision and will con-
   sult with a qualified medical professional if a medical judgment is
   involved. First level medical appeals will be [responded to in writ-
   ting to the member/ decided within thirty (30) business days for
   post-service claims and fifteen (15) days for pre-service claims] from
   the date the vendor received the first level appeal request.

   (a) If, because of reasons beyond the vendor’s control,
time more is needed to review the appeal, the vendor may extend the
time period up to an additional [fifteen (15)/ thirty (30) days]. The
vendor must notify the member prior to the expiration of the first
[fifteen- (15-) day/ twenty- (20-)] day period, explain the reason for the
delay, and request any additional information. If more information is
requested, the member has at least forty-five (45) days to provide the information to the vendor. The vendor then must decide the claim no later than [fifteen (15)] thirty (30) days after the additional information is supplied or after the period of time allowed to supply it ends, whichever is first. Written confirmation of the decision will be sent by the vendor within fifteen (15) business days.

(III) An expedited appeal of an adverse benefit determination may be requested when a decision is related to a pre-service claim for urgent care. Expedited appeals will be reviewed by the vendor by someone who was not involved in the original decision and will consult with a qualified medical professional if a medical judgment is involved. Expedited appeals will be responded to within seventy-two (72) hours after receiving a request for an expedited review with written confirmation of the decision to the member within three (3) working days of providing notification of the determination.

(IV) Second level appeals must be submitted in writing within sixty (60) days of the date of the first level appeal decision letter that upholds the original adverse benefit determination. Second level appeals should include any additional information or documentation to support the reason the member believes the first level appeal decision should be overturned. Second level appeals will be reviewed by the vendor by someone who was not involved in the original decision or first level appeal and will include consultation with a qualified medical professional if a medical judgment is involved. Second level medical appeals shall be responded to in writing to the member within thirty (30) days for post-service claims and within fifteen (15) days for pre-service claims from the date the vendor received the second level appeal request.

(a) If, because of reasons beyond the vendor’s control, more time is needed to review the appeal, the vendor may extend the time period up to an additional fifteen (15) days. The vendor must notify the member prior to the expiration of the first fifteen- (15-) day period, explain the reason for the delay, and request any additional information. If more information is requested, the member has at least forty-five (45) days to provide the information to the vendor. The vendor then must decide the claim no later than fifteen (15) days after the additional information is supplied or after the period of time allowed to supply it ends, whichever is first.

(V) For members with medical coverage through UMR—

(a) First and second level post-service, [first and second level post-service,] and concurrent claim appeals must be submitted in writing to—

| UMR Appeals |
| PO Box 400046 |
| San Antonio, TX 78229 |
| or by fax to (888) 615-6584 |

Anthem Blue Cross and Blue Shield
Attn: Grievance Department
PO Box 105568
Atlanta, Georgia 30348-5568
or by fax to (800) 859-3046

(b) First and second level post-service appeals must be sent in writing to—

| UMR Claims Appeal Unit |
| PO Box 30546 |
| Salt Lake City, UT 84130-0546 |
| or by fax to (877) 291-3248 |

(c)(b) Expedited [pre-service] appeals [must] may be [communicated] submitted by calling (800) 808-4424, ext. 15227, (877) 333-7488 or by submitting a written fax to (888) 615-6584, Attention: Appeals Unit (800) 368-3238.

(f) For members with medical coverage through Aetna—

(a) First and second level appeals must be submitted in writing to—

| Aetna Appeals Resolution Team |
| PO Box 14463 |
| Lexington, KY 40512 |
| or by fax to (859) 425-3379 |

(b) Expedited appeals must be communicated by calling (800) 245-0618 or by submitting a written fax to (859) 425-3379, Attention: Appeals Resolution Team.

C. The internal review process for adverse benefit determinations relating to pharmacy and the Pharmacy Lock-In Program consists of one (1) level of internal review provided by the pharmacy vendor.

(I) Pharmacy appeals. Pharmacy appeals and Pharmacy Lock-In Program appeals must identify the matter being appealed and shall include the member’s (and dependent’s, if applicable) name, the date the member attempted to fill the prescription, the prescribing physician’s name, the drug name and quantity, the cost of the prescription, if applicable, and any applicable reason(s) relevant to the appeal including: the reason(s) the member believes the claim should be paid, the reason(s) the member believes s/he should not be included in the Pharmacy Lock-In Program, and any other written documentation to support the member’s belief that the original decision should be overturned.

(II) All pharmacy appeals must be submitted in writing to—

Express Scripts
Attn: Clinical Appeals Department
PO Box 66588
St. Louis, MO 63116-6588
or by fax to (877) 852-4070

(III) All Pharmacy Lock-In Program appeals must be submitted in writing to—

Express Scripts
Drug Utilization Review Program
Mail Stop HQ3W03
One Express Way
St. Louis, MO 63121

(IV) Pharmacy appeals will be reviewed by someone who was not involved in the original decision and the reviewer will consult with a qualified medical professional if a medical judgment is involved. Pharmacy appeals will be responded to in writing to the member within sixty (60) days for post-service claims and thirty (30) days for pre-service claims from the date the vendor received the appeal request.

(V) The Pharmacy Benefit Manager will respond to Pharmacy Lock-In Program appeals in writing to the member within thirty (30) days from the date the Pharmacy Benefit Manager received the appeal request.

D. Members may seek external review only after they have exhausted all applicable levels of internal review or received a final internal adverse benefit determination.

(1) A claimant or authorized representative may file a written request for an external review within four (4) months after the date of receipt of a final internal adverse benefit determination.

(2) The claimant can submit an external review request in writing to—

HHS Federal Request
MAXIMUS Federal Services
3750 Monroe Ave., Suite 705
Pittsford, NY 14534
or by fax to (888) 866-6190
or to request a review online at http://www.externalappeal.com/
(III) The claimant may call the toll-free number (888) 866-6205 with any questions or concerns during the external review process and can submit additional written comments to the external reviewer at the mailing address above.

(IV) The external review decision will be made as expeditiously as possible and within forty-five (45) days after receipt of the request for the external review.

(V) A claimant may make a written or oral request for an expedited external review if the adverse benefit determination involves a medical condition of the claimant for which the timeframe for completion of a standard external review would seriously jeopardize the life or health of the claimant; or would jeopardize the claimant’s ability to regain maximum function; or if the final internal adverse benefit determination involves an admission, availability of care, continued stay, or health care item or service for which the claimant received services, but has not been discharged from a facility.

3. For all internal appeals of adverse benefit determinations, the plan or the vendor reviewing the appeal will provide the member, free of charge, with any new or additional evidence or rationale considered, relied upon, or generated by the plan or the vendor in connection with reviewing the claim or the appeal and will give the member an opportunity to respond to such new evidence or rationale before issuing a final internal adverse determination.

(5) In reviewing appeals, notwithstanding any other rule, the board and/or staff may grant any appeals when there is credible evidence to support appeal under the following guidelines. Decisions concerning eligibility for Medicare primary members may not be able to be granted pursuant to these guidelines if the decision is contrary to the rules controlling eligibility for Medicare Advantage plan as put forth by Centers for Medicare and Medicaid. Valid proof of eligibility must be included with the appeal if the enrollment request includes addition of dependent(s). Payment in full for all past and current premiums due for enrollment requests must be included with the appeal if it cannot be collected through payroll deduction.

(J) MCHCP may approve an appeal regarding plan changes retrospectively for subscribers who are new employees within thirty (30) days of election of coverage if no claims have been filed with the previous carrier, except that no changes will be considered for HSA Plan elections after the first MCHCP Health Savings Account contribution has been transmitted for deposit to the subscriber’s account. If a subscriber has her/his premium collected pre-tax by qualified payroll deduction through a cafeteria plan, changes may be approved if the reason given is allowed by the Missouri State Employees’ Cafeteria Plan; and

(K) Once per lifetime of the account, MCHCP may approve an appeal where a subscriber missed a deadline. MCHCP may only approve an appeal under this guideline if the appeal is received within sixty (60) days of the missed deadline. This guideline may not be used to approve an appeal of a voluntary cancellation or an appeal of a deadline that is statutorily mandated; and

[L] MCHCP may approve an appeal to change a subscriber’s medical plan vendor prospectively, once per lifetime of the account. This appeal guideline may not be used for a subscriber to change the type of medical plan design elected during open enrollment).


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 2—State Membership

EMERGENCY AMENDMENT

22 CSR 10-2.089 Pharmacy Employer Group Waiver Plan for Medicare Primary Members. The Missouri Consolidated Health Care Plan is amending section (1).

PURPOSE: This amendment updates the Medicare Part D coverage stage amounts and copayment amounts.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (employees, retirees, officers, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to members as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(1) The pharmacy benefit for Medicare primary non-active members is provided through a Pharmacy Employer Group Waiver Plan (EGWP) as regulated by the Centers for Medicare and Medicaid Services herein after referred to as the Medicare Prescription Drug Plan.

(F) The Medicare Prescription Drug Plan is comprised of a Medicare Part D prescription drug plan contracted by MCHCP and some non-Part D medications that are not normally covered by a Medicare Part D prescription drug plan. The requirements for the Medicare Part D prescription drug plan are as follows:

1. The Centers for Medicare and Medicaid Services regulates the Medicare Part D prescription drug program. The Medicare Prescription Drug Plan abides by those regulations;
2. Initial Coverage Stage. Until a member’s total yearly Part D prescription drug costs reach three thousand eight hundred
The Missouri Consolidated Health Care Plan is amending section (1).

PURPOSE: This amendment adds coinsurance dollar limits for the Health Savings Account (HSA) Plan, revises services covered at one hundred percent (100%), makes a technical correction to (1)(b)2.B., and renumbers as necessary.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (employees, retirees, officers, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unforeseen and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to members as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(I) The pharmacy benefit provides coverage for prescription drugs. Vitamin and nutrient coverage is limited to prenatal agents, therapeutic agents for specific deficiencies and conditions, and hematopoietic agents as prescribed by a provider to non-Medicare primary members.

(A) PPO 750 Plan and PPO 1250 Plan.

1. Network:

A. Preferred formulary generic drug copayment: Ten Dollars ($10) for up to a thirty-one- (31-) day supply; twenty dollars ($20) for up to a sixty- (60-) day supply; and thirty dollars ($30) for up to a ninety- (90-) day supply for a generic drug on the formulary.

B. Preferred formulary brand drug copayment: Forty dollars ($40) for up to a thirty-one- (31-) day supply; eighty dollars ($80) for up to a sixty- (60-) day supply; and one hundred twenty dollars ($120) for up to a ninety- (90-) day supply for a brand drug on the formulary.

C. Non-preferred formulary drug and approved excluded drug copayment: One hundred dollars ($100) for up to a thirty-one- (31-) day supply; two hundred dollars ($200) for up to a sixty- (60-) day supply; and three hundred dollars ($300) for up to a ninety- (90-) day supply for a drug not on the formulary.

D. Specialty drug copayment: Seventy-five dollars ($75) for up to a thirty-one- (31-) day supply for a specialty drug on the formulary.

E. Diabetic drug (as designated as such by the PBM) copayment: fifty percent (50%) of the applicable network copayment.

F. Home delivery programs.

(I) Maintenance prescriptions may be filled through the pharmacy benefit manager’s (PBM’s) home delivery program. A
member must choose how maintenance prescriptions will be filled by notifying the PBM of his/her decision to fill a maintenance prescription through home delivery or retail pharmacy.

(a) If the member chooses to fill his/her maintenance prescription at a retail pharmacy and the member does not notify the PBM of his/her decision, the first two (2) maintenance prescription orders may be filled by the retail pharmacy. After the first two (2) orders are filled at the retail pharmacy, the member must notify the PBM of his/her decision to continue to fill the maintenance prescription at the retail pharmacy. If a member does not make a decision after the first two (2) orders are filled at the retail pharmacy, s/he will be charged the full discounted cost of the drug until the PBM has been notified of the decision and the amount charged will not apply to the out-of-pocket maximum.

(b) Once a member makes his/her delivery decision, the member can modify the decision by contacting the PBM.

(II) Specialty drugs are covered only through the specialty home delivery network for up to a thirty-one-(31-) day supply unless the PBM has determined that the specialty drug is eligible for up to a ninety-(90-) day supply. All specialty prescriptions must be filled through the PBM’s specialty pharmacy, unless the prescription is identified by the PBM as emergent. The first fill of a specialty prescription identified to be emergent, may be filled through a retail pharmacy.

(a) Specialty split-fill program—The specialty split-fill program applies to select specialty drugs as determined by the PBM. For the first three (3) months, members will be shipped a fifteen-(15-) day supply and charged a prorated copayment. If the member is able to continue with the medication, the remaining supply will be shipped and the member will be charged the remaining portion of the copayment. Starting with the fourth month, an up to thirty-one-(31-) day supply will be shipped if the member continues on treatment.

(III) Prescriptions filled through home delivery programs have the following copayments:

(a) Preferred formulary generic drug copayments: Ten dollars ($10) for up to a thirty-one-(31-) day supply; twenty dollars ($20) for up to a sixty-(60-) day supply; and twenty-five dollars ($25) for up to a ninety-(90-) day supply for a generic drug on the formulary;

(b) Preferred formulary brand drug copayments: Forty dollars ($40) for up to a thirty-one-(31-) day supply; eighty dollars ($80) for up to a sixty-(60-) day supply; and one hundred dollars ($100) for up to a ninety-(90-) day supply for a brand drug on the formulary;

(c) Non-preferred formulary drug and approved excluded drug copayments: One hundred dollars ($100) for up to a thirty-one-(31-) day supply; two hundred dollars ($200) for up to a sixty-(60-) day supply; and two hundred fifty dollars ($250) for up to a ninety-(90-) day supply for a drug not on the formulary;

(d) Specialty drug copayment: Seventy-five dollars ($75) for up to a thirty-one-(31-) day supply; one hundred fifty dollars ($150) for up to sixty-(60-) day supply; and two hundred twenty-five dollars ($225) for up to ninety-(90-) day supply for a specialty drug on the formulary;

E. Diabetic drug (as designated as such by the PBM) copayment: fifty percent (50%) of the applicable network copayment;

F. Only one (1) copayment is charged if a combination of different manufactured dosage amounts must be dispensed in order to fill a prescribed single dosage amount;

I. The copayment for a compound drug is based on the primary drug in the compound. The primary drug in a compound is the most expensive prescription drug in the mix. If any ingredient in the compound is excluded by the plan, the compound will be denied;

J. If the copayment amount is more than the cost of the drug, the member is only responsible for the cost of the drug;

K. If the physician allows for generic substitution and the member chooses a brand-name drug, the member is responsible for the generic copayment and the cost difference between the brand-name and generic drug which shall not apply to the out-of-pocket maximum;

L. Preferred select brand drugs, as determined by the PBM: Ten dollars ($10) for up to a thirty-one-(31-) day supply; twenty dollars ($20) for up to a sixty-(60-) day supply; and twenty-five dollars ($25) for up to a ninety-(90-) day supply; and

M. Prescription drugs and prescribed over-the-counter drugs as recommended by the U.S. Preventive Services Task Force (categories A and B) and, for women, by the Health Resources and Services Administration are covered at one hundred percent (100%) when filled at a network pharmacy. The following are also covered at one hundred percent (100%) when filled at a network pharmacy:

1. Vaccine recommended by the Advisory Committee on Immunization Practices of the Centers for Disease Control and Prevention;

2. [(III) Joniper Tamoxifen, generic Raloxifene, and brand Softamox for prevention of breast cancer;]

3. [(IV)(II) Prescribed preferred diabetic test strips and lancets; and]

4. [(V)(III) One (1) preferred glucometer.]

2. Non-network: If a member chooses to use a non-network pharmacy for non-specialty prescriptions, s/he will be required to pay the full cost of the prescription and then file a claim with the PBM. The PBM will reimburse the cost of the drug based on the network discounted amount as determined by the PBM, less the applicable network copayment.


A. Network and non-network out-of-pocket maximums are separate.

B. The family out-of-pocket maximum is an aggregate of applicable charges received by all covered family members of the plan. Any combination of covered family member applicable charges may be used to meet the family out-of-pocket maximum. Applicable charges received by one (1) family member may only meet the individual out-of-pocket maximum amount.

C. Network individual—four thousand one hundred fifty dollars ($4,150).

D. Network family—eight thousand three hundred dollars ($8,300).

E. Non-network—no maximum.

(B) Health Savings Account (HSA) Plan Prescription Drug Coverage. Medical and pharmacy expenses are combined to apply toward the appropriate network or non-network deductible and out-of-pocket maximum specified in 22 CSR 10-2.053.

1. Network:

A. Preferred formulary generic drug: Ten percent (10%) coinsurance up to fifty dollars ($50) per thirty-one-(31-) day supply after deductible has been met for a generic drug on the formulary;

B. Preferred formulary brand drug: Twenty percent (20%) coinsurance up to one hundred dollars ($100) per thirty-one-(31-) day supply after deductible has been met for a brand drug on the formulary;

C. Non-preferred formulary drug and approved excluded drug: Forty percent (40%) coinsurance after deductible has been met;

D. Diabetic drug (as designated as such by the PBM) coinsurance: fifty percent (50%) of the applicable network coinsurance after deductible has been met, not to exceed:

1. Twenty-five dollars ($25) per thirty-one-(31-) day supply for generic drugs;

2. Fifty dollars ($50) per thirty-one-(31-) day supply for preferred formulary brand drug; and

3. One hundred dollars ($100) per thirty-one-(31-) day supply for non-preferred formulary drug;

E. Home delivery programs.
(I) Maintenance prescriptions may be filled through the PBM’s home delivery program. A member must choose how maintenance prescriptions will be filled by notifying the PBM of his/her decision to fill a maintenance prescription through home delivery or retail pharmacy.

(a) If the member chooses to fill his/her maintenance prescription at a retail pharmacy and the member does not notify the PBM of his/her decision, the first two (2) maintenance prescription orders may be filled at the retail pharmacy. After the first two (2) orders are filled at the retail pharmacy, the member must notify the PBM of his/her decision to continue to fill the maintenance prescription at the retail pharmacy. If a member does not make a decision after the first two (2) orders are filled at the retail pharmacy, s/he will be charged the full discounted cost of the drug until the PBM has been notified of the decision.

(b) Once a member makes his/her delivery decision, the member can modify the decision by contacting the PBM.

(II) Specialty drugs are covered only through the specialty home delivery network for up to a thirty-one- (31-) day supply unless the PBM has determined that the specialty drug is eligible for up to a ninety- (90-) day supply. All specialty prescriptions must be filled through the PBM’s specialty pharmacy, unless the prescription is identified by the PBM as emergent. The first fill of a specialty prescription identified to be emergent, may be filled through a retail pharmacy.

(a) Specialty split-fill program—The specialty split-fill program applies to select specialty drugs as determined by the PBM. For the first three (3) months, members will be shipped a fifteen- (15-) day supply. If the member is able to continue with the medication, the remaining supply will be shipped. Starting with the fourth month, an up to thirty-one- (31-) day supply will be shipped if the member continues on treatment.

F. Prescription drugs and prescribed over-the-counter drugs as recommended by the U.S. Preventive Services Task Force (categories A and B) and, for women, by the Health Resources and Services Administration are covered at one hundred percent (100%) when filled at a network pharmacy. [The following are also covered at one hundred percent (100%) when filled at a network pharmacy:]

[I]G. Vaccines and administration as recommended by the Advisory Committee on Immunization Practices of the Centers for Disease Control and Prevention are covered at one hundred percent (100%) when filled at a network pharmacy;

[II]H. Diabetic drug (as designated as such by the PBM) coinsurance: fifty percent (50%) of the applicable non-network coinsurance after deductible has been met.


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 2—State Membership

EMERGENCY AMENDMENT

22 CSR 10-2.110 General Foster Parent Membership Provisions. The Missouri Consolidated Health Care Plan is amending sections (3), (5), and (14).

PURPOSE: This amendment revises default enrollment procedures, clarifies disabled dependent eligibility, reporting of other health coverage, and renumbers as necessary.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (employees, retirees, officers, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to members in one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(3) Enrollment Procedures.
EMERGENCY RULES

(C) An eligible foster parent may elect or change coverage for himself/herself and/or for his/her spouse/child(ren) if one (1) of the following occurs:

1. Occurrence of a life event, which includes marriage, birth, adoption, and placement of child(ren). A special enrollment period of thirty-one (31) days shall be available beginning with the date of the life event. It is the eligible foster parent’s responsibility to notify MCHCP of the life event:
   A. If paternity is necessary to establish the life event and was not established at birth, the date that paternity is established shall be the date of the life event; or
   B. Employer-sponsored group coverage loss. An eligible foster parent or his/her spouse/child(ren) may enroll within sixty (60) days due to an involuntary loss of employer-sponsored coverage under one (1) of the following circumstances:
      A. Employer-sponsored medical, dental, or vision coverage terminates;
      B. Eligibility for employer-sponsored coverage ends;
      C. Employer contributions toward the premiums end; or
      D. Consolidated Omnibus Budget Reconciliation Act (COBRA) coverage ends;
   C. If an eligible foster parent or his/her spouse/child(ren) loses MO HealthNet or Medicaid status, s/he may enroll in an MCHCP plan within sixty (60) days of the date of loss; or
   D. If an eligible foster parent or eligible foster parent’s spouse receives a court order stating s/he is responsible for covering a child, the eligible foster parent may enroll the child in an MCHCP plan within sixty (60) days of the court order; or
   E. 5. Default Enrollment
      A. If an eligible foster parent is enrolled in the PPO [300 or] 750, PPO /[600] 1250, or HSA Plan and does not complete enrollment during the open enrollment period, the foster parent and his/her dependents will be enrolled in the same plan at the same level of coverage [in the PPO 1250 Plan provided through the vendor] the foster parent is enrolled in, effective the first day of the next calendar year; or
      B. If an eligible foster parent is enrolled in the Health Savings Account (HSA) Plan and does not complete enrollment during the open enrollment period, the foster parent and his/her dependents will be enrolled in the same level of coverage in the HSA Plan provided through the vendor the foster parent is enrolled in, effective the first day of the next calendar year;
      C/B. If an eligible foster parent is enrolled in dental and/or vision coverage and does not complete open enrollment to cancel coverage or change the current level of coverage during the open enrollment period, the foster parent and his/her dependents will be enrolled at the same level of coverage in the same plan(s), effective the first day of the next calendar year; or
   6. If an eligible foster parent submits an Open Enrollment Worksheet or an Enroll/Change/Cancel form that is incomplete or contains obvious errors, MCHCP will notify the foster parent of such by mail, phone, or secure message. The foster parent must submit a corrected form to MCHCP by the date enrollment was originally due to MCHCP or ten (10) business days from the date MCHCP notifies the foster parent, whichever is later.

(5) Proof of Eligibility. Proof of eligibility documentation is required for all dependents and subscribers, as necessary. Enrollment is not complete until proof of eligibility is received by MCHCP. A subscriber must include his/her MCHCPid or Social Security number on the documentation. If proof of eligibility is not received, MCHCP will send a letter requesting it from the subscriber. Except for open enrollment, documentation must be received within thirty-one (31) days of the date MCHCP processed the enrollment, or coverage will not take effect for those individuals whose proof of eligibility was not received. MCHCP reserves the right to request that such proof of eligibility be provided at any time upon request. If such proof is not received or is unacceptable as determined by MCHCP, coverage will terminate or never take effect. If enrolling during open enrollment, proof of eligibility must be received by November 20, or coverage will not take effect the following January 1 for those individuals whose proof of eligibility was not received. If invalid proof of eligibility is received, the subscriber is allowed an additional ten (10) days from the initial due date to submit valid proof of eligibility.

(E) Disabled Dependent

1. An [newly] eligible foster parent may enroll his/her permanently disabled child when first eligible or an enrolled permanently disabled dependent turning age twenty-six (26) years, may continue coverage beyond age twenty-six (26) years, provided the following documentation is submitted to the plan prior to the end of the month of the dependent’s twenty-sixth birthday for the enrolled permanently disabled dependent or within thirty-one (31) days of enrollment of a new foster parent and his/her the permanently disabled child:
   A. Evidence from the Social Security Administration (SSA) that the permanently disabled dependent or child was entitled to and receiving disability benefits prior to turning age twenty-six (26) years; and
   B. A benefit verification letter dated within the last twelve (12) months from the SSA confirming the child is still considered disabled.
   2. If a disabled dependent over the age of twenty-six (26) years is determined to be no longer disabled by the SSA, coverage will terminate the last day of the month in which the disability ends or never take effect for new enrollment requests.
   3. Once the disabled child’s coverage is cancelled or terminated, s/he will not be able to enroll at a later date.

(14) Members are required to disclose to the claims administrator whether they have other health coverage and, if so, information about the coverage. A member may submit other coverage information to the claims administrator by phone, fax, mail, or online. Dependent claims will be denied until the information is received. Once the information is received, claims will be reprocessed subject to all applicable rules.


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—Missouri Consolidated Health Care Plan
Division 10—Health Care Plan
Chapter 3—Public Entity Membership

EMERGENCY AMENDMENT

22 CSR 10-3.020 General Membership Provisions. The Missouri Consolidated Health Care Plan is amending sections (5) and (13).

PURPOSE: This amendment clarifies disabled dependent eligibility and reporting of other health coverage.

EMERGENCY STATEMENT: This emergency amendment must be in
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Missouri Register

EMERGENCY RESCISSION

22 CSR 10-3.045 Plan Utilization Review Policy. This rule established the policy of the board of trustees in regard to the Plan Utilization Review Policy of the Missouri Consolidated Health Care Plan.

PURPOSE: This rule is being rescinded and readopted to reflect changes due to a new third party administrator.

EMERGENCY STATEMENT: This emergency rescission must be in place by January 1, 2020 in accordance with the new plan year. Therefore, this emergency rescission is necessary to serve a compelling governmental interest of protecting members (public entity employee members, retirees, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this rescission be filed as an emergency rescission to maintain the integrity of the current health care plan. This emergency rescission fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to public entity employee members, retirees, and their families as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency rescission reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed rescission, which covers the same material, is published in this issue of the Missouri Register. This emergency rescission complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency rescission was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN

Chapter 10—Health Care Plan

Chapter 3—Public Entity Membership


PUBLIC COST: This emergency rescission will not cost state agencies or political subdivisions more than five hundred dollars ($500)
in the aggregate.

PRIVATE COST: This emergency rescission will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 3—Public Entity Membership

EMERGENCY RULE

22 CSR 10-3.045 Plan Utilization Review Policy

PURPOSE: This rule establishes the policy of the board of trustees in regard to the Plan Utilization Review Policy of the Missouri Consolidated Health Care Plan.

EMERGENCY STATEMENT: This emergency rule must be in place by January 1, 2020 in accordance with the new plan year. Therefore, this emergency rule is necessary to serve a compelling governmental interest of protecting members (public entity employee members, retirees, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this rule be filed as an emergency rule to maintain the integrity of the current health care plan. This emergency rule fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to public entity employee members, retirees, and their families as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency rule reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed rule, which covers the same material, is published in this issue of the Missouri Register. This emergency rule complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency rule was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(1) Clinical Management—Certain benefits are subject to a utilization review (UR) program. The program has the following components:

(A) Preauthorization—The claims administrator must authorize some services in advance. Preauthorization is to determine if the procedure or treatment is medically necessary. The claims administrator will determine what procedures or treatments are subject to preauthorization. Without preauthorization, any claim that requires preauthorization will be denied for payment. Members who have another primary carrier, or who are enrolled in the Medicare Advantage Plan are not subject to this provision except for those services that are not covered by the other primary carrier, but are otherwise subject to preauthorization under this rule. Preauthorizations found to have a material misrepresentation or intentional or negligent omission about the person’s health condition or the cause of the condition may be rescinded.

1. A list of medical services for which preauthorization is required may be obtained at any time from the claims administrator.

2. The following pharmacy services included in the prescription drug plan for non-Medicare primary members are subject to preauthorization:

A. Second-step therapy medications that skip the first-step medication trial;
B. Specialty medications;
C. Medications that may be prescribed for several conditions, including some for which treatment is not medically necessary;
D. Medication refill requests that are before the time allowed for refill;
E. Medications that exceed drug quantity and day supply limitations; and
F. Medications with costs exceeding nine thousand nine hundred ninety-nine dollars and ninety-nine cents ($9,999.99) at retail or the mail order pharmacy and one hundred forty-nine dollars and ninety-nine cents ($149.99) for compound medications at retail or the mail order pharmacy.

3. Preauthorization timeframes.

A. A benefit determination for non-urgent preauthorization requests will be made within thirty-six (36) hours, which will include one (1) business day of the receipt of the request. If the information necessary to make a benefit determination is not received, the claims administrator will notify the member and provider of any necessary extension. The provider will be given forty-five (45) calendar days from receipt of the extension notice to respond with additional information. Once the information is received or the forty-five (45) days have elapsed, a determination will be made within thirty-six (36) hours which will include one (1) business day.

B. A benefit determination for urgent preauthorization requests will be made as soon as possible based on the clinical situation, but in no case later than one (1) business day of the receipt of all necessary information;

(B) Concurrent Review—The claims administrator will monitor the medical necessity of an inpatient admission to certify the necessity of the continued stay in the hospital. Members who have another primary carrier, including Medicare, are not subject to this provision;

(C) Retrospective Review—Reviews to determine coverage after services have been provided to a member. The retrospective review is not limited to an evaluation of medical necessity, reimbursement levels, accuracy and adequacy of documentation or coding, or settling of payment. The claim administrator shall have the authority to correct payment errors when identified under retrospective review;

(D) Pre-determination—Determination of coverage by the claims administrator prior to services being provided. A provider may voluntarily request a pre-determination. A pre-determination informs the provider of whether, and under which circumstances, a procedure or service is generally a covered benefit under the plan. A pre-determination that a procedure or service may be covered under the plan does not guarantee payment; and

(E) Case Management—A voluntary process to assess, coordinate, and evaluate options and services of members with catastrophic and complex illnesses. A case manager will help members understand what to expect during the course of treatment, help establish collaborative goals, complete assessments to determine needs, interface with providers, and negotiate care. Members are identified for case management through claim information, length of hospital stay, or by referral. The case manager will dismiss the member from case management once the case manager determines that objectives have been met.

PUBLIC COST: This emergency rule will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency rule will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 3—Public Entity Membership

EMERGENCY AMENDMENT

22 CSR 10-3.055 Health Savings Account Plan Benefit Provisions and Covered Charges. The Missouri Consolidated Health Care Plan is amending sections (3), (13), (14), (18), and adding section (9).

PURPOSE: This amendment revises the Health Savings Account (HSA) Plan individual family member out-of-pocket maximum, adds one hundred percent (100%) coverage after deductible is met for sterilization procedure for men, revises maximum plan payment, timely filing timeframe, services performed in another country, and renumbers as necessary.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (public entity employee members, retirees, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to public entity employee members, retirees, and their families as one method of protecting the MCHCP trust fund from more costly claims and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(3) Out-of-pocket maximum.

(A) The family out-of-pocket maximum applies when two (2) or more family members are covered. The family out-of-pocket maximum must be met before the plan begins to pay one hundred percent (100%) of all covered charges for any covered family member. Out-of-pocket maximums are per calendar year, as follows:

1. Network out-of-pocket maximum for individual—four thousand nine hundred fifty dollars ($4,950);
2. Network out-of-pocket maximum for family—nine thousand nine hundred dollars ($9,900).

Any individual family member need only incur a maximum of [seven thousand nine hundred dollars ($7,900)] eight thousand one hundred fifty dollars ($8,150) before the plan begins paying one hundred percent (100%) of covered charges for that individual;

3. Non-network out-of-pocket maximum for individual—nine thousand nine hundred dollars ($9,900); and

(9) Sterilization procedure for men is paid at one hundred percent (100%) when provided by a network provider after deductible is met.

//9//10 Newborn’s claims will be subject to deductible and coinsurance.

//10//11 Each subscriber will have access to payment information of the family unit only when authorization is granted by the adult covered dependent(s).

//11//12 Expenses toward the deductible and out-of-pocket maximum will be transferred if the member changes medical plans or continues enrollment under another subscriber’s plan within the same plan year.

//12//13 Maximum plan payment—Non-network medical claims that are not otherwise subject to a contractual discount arrangement are processed at one hundred ten percent (110%) of Medicare reimbursement for non-network professional claims and following the claim administrator’s standard practice for non-network facility claims. Members may be held liable for the amount of the fee above the allowed amount.

//13//14 Any claim must be initially submitted within twelve (12) months following the date of service, unless other specified in the network provider contract. The plan reserves the right to deny claims not timely filed. A provider initiated correction to the originally filed claim must be submitted within the timeframe agreed in the provider contract, but not to exceed three hundred sixty-five (365) days from adjudication of the originally filed claim. Any claims reprocessed as primary based on action taken by Medicare or Medicaid must be initiated within three (3) years of the claim being incurred.

//14//15 For a member who is an inpatient on the last calendar day of a plan year and remains an inpatient into the next plan year, the prior plan year’s applicable deductible and/or coinsurance amounts will apply to the in-hospital facility and related ancillary charges until the member is discharged.

//15//16 A subscriber does not qualify for the HSA Plan if s/he is claimed as a dependent on another person’s tax return or, except for the plans listed in section (16) of this rule, is covered under or enrolled in any other health plan that is not a high deductible health plan, including, but not limited to, the following types of insurance plans or programs:

(A) Medicare (unless Medicare is secondary coverage to MCHCP);
(B) TRICARE;
(C) A health care flexible spending account (FSA), with the exception of participation in the premium-only, limited-purpose health FSA, and dependent care section;
(D) Health reimbursement account (HRA); or
(E) If the member has received medical benefits from The Department of Veterans Affairs (VA) at any time during the previous three (3) months, unless the medical benefits received consist solely of disregarded coverage or preventive care.
A subscriber may qualify for this plan even if s/he is covered by any of the following:
(A) Drug discount card;
(B) Accident insurance;
(C) Disability insurance;
(D) Dental insurance;
(E) Vision insurance; or
(F) Long-term care insurance.

Services performed in a country other than the United States may be covered if the service is included in 22 CSR 10-3.057. Emergency and urgent care services are covered as a network benefit. All other non-emergency services are covered as a non-network benefit determined by the claims administrator. If the service is provided by a non-network provider, the member may be required to provide payment to the provider and then file a claim for reimbursement subject to timely filing limits.


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 3—Public Entity Membership

EMERGENCY AMENDMENT

22 CSR 10-3.057 Medical Plan Benefit Provisions and Covered Charges. The Missouri Consolidated Health Care Plan is amending sections (2) and (3).

PURPOSE: This amendment revises the medical plan benefits for transition of care, allergy testing and immunotherapy, bariatric surgery, bone growth stimulators, cardiac rehabilitation, chelation therapy, chiropractic services, cochlear implant, dental care, emergency room services, genetic counseling, hearing aids, hospice care and palliative services, hospital, injections, maternity coverage, nutrition therapy, orthognathic or jaw surgery, orthotics, and renum bers as necessary.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (public entity employee members, retirees, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to public entity employee members, retirees, and their families as one method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(1) Transition of Care. A transition of care option is available for members using a hospital or dialysis facility that loses network status during the plan year. A subscriber and his/her dependents may apply for additional days beyond the ninety-(90-) day transition of care to continue receiving network benefits with that hospital or dialysis facility. The request for consideration must be submitted to the medical plan within forty-five (45) days of the last day the hospital or dialysis facility was a contracted network provider, to be eligible for transition of care benefits. A subscriber and his/her dependents may apply for additional days beyond the ninety-(90-) day transition if care is related to a moderate or high risk pregnancy, if care is during a member’s second or third trimester of pregnancy, or up to eight (8) weeks postpartum. The subscriber and his/her dependents must apply for additional transition of care days prior to the end of the initial ninety-(90-) day transition of care period. Most routine services, treatment for stable conditions, minor illnesses, and elective surgeries will not be covered by transition of care benefits. The rate of payment during the transitional period shall be the fee paid prior to leaving the network. Benefits eligible for transition of care include:
(A) Upcoming surgery or prospective transplant;
(B) Services for women in their second or third trimester of pregnancy or up to eight (8) weeks postpartum;
(C) Services for women who have been diagnosed as potentially having a moderate- or high-risk pregnancy;
(D) Home nursing care;
(E) Radiation therapy;
(F) Dialysis;
(G) Durable medical equipment;
(H) Cancer treatment;
(I) Clinical trials;
(J) Physical, speech, or occupational therapy;
(K) Hospice care;
(L) Bariatric surgery, and follow-up per criteria covered under the plan;
(M) Inpatient hospitalization at the time of the network change;
(N) Mental health services; or
(O) Related follow-up services within three (3) months of an acute injury or surgery.

(2) Transition of Care. A transition of care option is available for members who seek to continue to remain under the care of an non-network provider who was treating them prior to the provider losing network status. A subscriber and his/her dependents may
request to continue receiving care at the network benefit level. If approved, the member will be eligible to continue care with the current non-network provider at the network benefit level for a period of time until it is medically appropriate for the member to transfer care to a network provider. The rate of payment during the transitional period shall be the fee paid prior to leaving the network. The following benefits are eligible for transition of care as determined by the claims administrator:

(A) Upcoming surgery or prospective transplant;
(B) Services for women in their third trimester of pregnancy;
(C) Radiation therapy;
(D) Dialysis;
(E) Cancer treatment;
(F) Physical, speech, or occupational therapy;
(G) Hospice care;
(H) Inpatient hospitalization at the time of the network change;

(1) Mental health services.

(3) Covered Charges Applicable to the PPO 750 Plan, PPO 1250, and HSA Plan.

[(ID Services in a country other than the United States. Emergency room and urgent care medical services are covered at the network benefit. All other non-emergency services are covered at the non-network benefit.)

[(EI)(D) Plan benefits for the PPO 750 Plan, PPO 1250, and HSA Plan are as follows:

1. Allergy Testing and Immunotherapy. Allergy testing and allergy immunotherapy are considered medically necessary for members with clinically significant allergic symptoms. The following tests and treatments are covered:

   A. Epicutaneous (scratch, prick, or puncture) when Immunoglobulin E- (IgE-) mediated reactions occur to any of the following:
      (1) Foods;
      (2) Hymenoptera venom (stinging insects);
      (3) Inhalants; or
      (4) Specific drugs (penicillins and macromolecular agents);

   B. Intradermal (Intracutaneous) when IgE-mediated reactions occur to any of the following:
      (1) Foods;
      (2) Hymenoptera venom (stinging insects);
      (3) Inhalants; or
      (4) Specific drugs (penicillins and macromolecular agents);

   C. Skin or Serial Endpoint Titration (SET), also known as intradermal dilutional testing (IDT), for determining the starting dose for immunotherapy for members highly allergic to any of the following:
      (1) Hymenoptera venom (stinging insects); or
      (2) Inhalants;

   D. Skin Patch Testing: for diagnosing contact allergic dermatitis;

   E. Photo Patch Testing: for diagnosing photo-allergy (such as photo-allergic contact dermatitis);

   F. Photo Tests: for evaluating photo-sensitivity disorders;

   G. Bronchial Challenge Test: for testing with methacholine, histamine, or antigens in defining asthma or airway hyperactivity when either of the following conditions is met:
      (1) Bronchial challenge test is being used to identify new allergens for which skin or blood testing has not been validated; or
      (2) Skin testing is unreliable;

   H. Exercise Challenge Testing for exercise-induced bronchospasm;

   I. Ingestion (Oral) Challenge Test for any of the following:
      (1) Food or other substances; or
      (2) Drugs when all of the following are met:
         (a) History of allergy to a particular drug;
         (b) There is no effective alternative drug; and
         (c) Treatment with that drug class is essential;

   J. In Vitro IgE Antibody Tests (RAST, MAST, FAST, ELISA, ImmunoCAP) are covered for the following:
      (1) Allergic broncho-pulmonary aspergillosis (ABPA) and certain parasitic diseases;
      (2) Food allergy;
      (3) Hymenoptera venom allergy (stinging insects);
      (4) Inhalant allergy; or
      (5) Specific drugs;

   K. Total Serum IgE for diagnostic evaluation in members with known or suspected ABPA and/or hyper IgE syndrome;

   L. Lymphocyte transformation tests such as lymphocyte mitogen response test, PHE stimulation test, or lymphocyte antigen response assay are covered for evaluation of persons with any of the following suspected conditions:
      (1) Sensitivity to beryllium;
      (2) Congenital or acquired immunodeficiency diseases affecting cell-mediated immunity, such as severe combined immunodeficiency, common variable immunodeficiency, X-linked immunodeficiency with hyper IgM, Nijmegen breakage syndrome, reticular dysgenesis, DiGeorge syndrome, Nezelof syndrome, Wiscott-Aldrich syndrome, ataxia telangiectasia, and chronic mucocutaneous candidiasis;
      (3) Thymoma; and

   (IV) To predict allograft compatibility in the transplant setting;

   M. Allergy retesting: routine allergy retesting is not considered medically necessary;

   N. Allergy immunotherapy is covered for the treatment of any of the following IgE-mediated allergies:
      (1) Allergic (extrinsic) asthma;
      (2) Dust mite atopic dermatitis;
      (3) Hymenoptera (bees, hornets, wasps, fire ants) sensitive individuals;
      (4) Mold-induced allergic rhinitis;
      (5) Perennial rhinitis;
      (6) Seasonal allergic rhinitis or conjunctivitis when one (1) of the following conditions are met:
         (a) Member has symptoms of allergic rhinitis or asthma after natural exposure to the allergen;
         (b) Member has a life-threatening allergy to insect stings; or
         (c) Member has skin test or serologic evidence of IgE mediated antibody to a potent extract of the allergen;

   (VII) Avoidance or pharmacologic therapy cannot control allergic symptoms or member has unacceptable side effects with pharmacologic therapy;

   O. Other treatments: the following other treatments are covered:
      (1) Rapid, rush, cluster, or acute desensitization for members with any of the following conditions:
         (a) IgE antibodies to a particular drug that cannot be treated effectively with alternative medications;
         (b) Insect sting (e.g., wasps, hornets, bees, fire ants) hypersensitivity (hymenoptera); or
         (c) Members with moderate to severe allergic rhinitis who need treatment during or immediately before the season of the affecting allergy;

      (II) Rapid desensitization is considered experimental and investigational for other indications;
P. Epinephrine kits, to prevent anaphylactic shock for members who have had life-threatening reactions to insect stings, foods, drugs, or other allergens; have severe asthma or if needed during immunotherapy;

2. Ambulance service. The following ambulance transportation services are covered:
   A. By ground to the nearest appropriate facility when other means of transportation would be contraindicated;
   B. By air to the nearest appropriate facility when the member’s medical condition is such that transportation by either basic or advanced life support ground ambulance is not appropriate or contraindicated;

3. Applied Behavior Analysis (ABA) for Autism;
4. Bariatric surgery. Bariatric surgery is covered when all of the following requirements have been met:
   [A. The surgery is performed at a facility accredited by the Metabolic and Bariatric Surgery Accreditation and Quality Improvement Program (MBSAQIP) for the billed procedure;]
   B. The following open or laparoscopic bariatric surgery procedures are covered:
      (I) Roux-en-Y gastric bypass;
      (II) Sleeve gastrectomy;
      (III) Biliopancreatic diversion with duodenal switch for individuals with a body mass index (BMI) greater than 50;
      (IV) Adjustable silicone gastric banding and adjustments of a silicone gastric banding to control the rate of weight loss and/or treat symptoms secondary to gastric restriction following an adjustable silicone gastric banding procedure;
      (V) Surgical reversal of bariatric surgery when complications of the original surgery (e.g., stricture, pouch dilatation, erosion, or band slippage) cause abdominal pain, inability to eat or drink, or cause vomiting of prescribed meals;
   [VI] Revision of a previous bariatric surgical procedure or conversion to another procedure due to inadequate weight loss when one (1) of the following specific criteria has been met:
      (a) There is evidence of full compliance with the previously prescribed post-operative dietary and exercise program; or
      (b) There is documented clinical testing demonstrating technical failure of the original bariatric surgical procedure which caused the individual to fail achieving adequate weight loss of at least fifty percent (50%) of excess body weight or failure to achieve body weight to within thirty percent (30%) of ideal body weight at least two (2) years following the original surgery;
   C. All of the following criteria have been met:
      (I) The member is eighteen (18) years or older or has reached full skeletal growth, and has evidence of one (1) of the following:
         (a) BMI greater than forty (40); or
         (b) BMI between thirty-five (35) and thirty-nine and nine tenths (39.9) and one (1) or more of the following:
            I. Type II diabetes;
            II. Cardiovascular disease such as stroke, myocardial infarction, stable or unstable angina pectoris, hypertension, or coronary artery bypass; or
            III. Life-threatening cardiopulmonary problems such as severe sleep apnea, Pickwickian syndrome, or obesity-related cardiomyopathy; and
      (II) Demonstration that dietary attempts at weight control have been ineffective through completion of a structured diet program. Commercial weight loss programs are acceptable if completed under the direction of a provider or registered dietitian and documentation of participation is available for review. One (1) structured diet program for six (6) consecutive months or two (2) structured diet programs for three (3) consecutive months each within a two (2)-year period prior to the request for the surgical treatment of morbid obesity are sufficient. Provider-supervised programs consisting exclusively of pharmacological management are not sufficient; and
      (III) A thorough multidisciplinary evaluation within the previous twelve (12) months, which include all of the following:
         (a) An evaluation by a bariatric surgeon recommending surgical treatment, including a description of the proposed procedure and all of the associated current procedural terminology codes;
         (b) A separate medical evaluation from a provider other than the surgeon recommending surgery that includes a medical clearance for bariatric surgery;
         (c) Completion of a psychological examination from a mental health provider evaluating the member’s readiness and fitness for surgery and the necessary post-operative lifestyle changes. After the evaluation, the mental health provider must provide clearance for bariatric surgery; and
         (d) A nutritional evaluation by a provider or registered dietitian;

5. Blood storage. Storage of whole blood, blood plasma, and blood products is covered in conjunction with medical treatment that requires immediate blood transfusion support;
6. Bone Growth Stimulators. Implantable bone growth stimulators are covered as an outpatient surgery benefit. The following nonimplantable bone growth stimulators are covered as a durable medical equipment benefit:
   [A. Ultrasound osteogenesis stimulator (e.g., the Sonic Accelerated Fracture Healing System (SAFHS)) to accelerate healing of fresh fractures, fusions, or delayed unions at either of the following high-risk sites:
      (I) Fresh fractures, fusions, or delayed unions of the shaft (diaphysis) of the tibia that are open or segmental; or
      (II) Fresh fractures, fusions, or delayed unions of the scaphoid (carpal navicular);
   B. Ultrasound osteogenesis stimulator for non-unions, failed arthrodesis, and congenital pseudarthrosis (pseudarthrosis) of the appendicular skeleton if there has been no progression of healing for three (3) or more months despite appropriate fracture care; or
   C. Direct current electrical bone-growth stimulator is covered for the following indications:
      (I) Delayed unions of fractures or failed arthrodesis at high-risk sites (i.e., open or segmental tibial fractures, carpal navicular fractures);
      (II) Non-unions, failed fusions, and congenital pseudarthrosis where there is no evidence of progression of healing for three (3) or more months despite appropriate fracture care; or
      (III) Members who are at high risk for spinal fusion failure when any of the following criteria is met:
         (a) A multiple-level fusion entailing three (3) or more vertebrae (e.g., L3 to L5, L4 to S1, etc.);
         (b) Grade II or worse spondylolisthesis; or
         (c) One (1) or more failed fusions;]
7. Contraception and Sterilization. All Food and Drug Administration- (FDA-) approved contraceptive methods, sterilization procedures, and patient education and counseling for all women with reproductive capacity;
8. Cardiac rehabilitation. An electrocardiographically-monitored program of outpatient cardiac rehabilitation (Phase II) is covered for specific criteria when it is individually prescribed by a provider and a formal exercise stress test is completed following the event and prior to the initiation of
the program. Cardiac rehabilitation is covered for members who meet one (1) of the following criteria:

[A. Acute myocardial infarction (MI) (heart attack in the last twelve (12) months);
B. Coronary artery bypass grafting (CABG);
C. Stable angina pectoris;
D. Percutaneous coronary vessel remodeling;
E. Valve replacement or repair;
F. Heart transplant;
G. Coronary artery disease (CAD) associated with chronic stable angina that has failed to respond adequately to pharmacotherapy and is interfering with the ability to perform age-related activities of daily living and/or impairing functional abilities; or
H. Heart failure that has failed to respond adequately to pharmacotherapy and is interfering with the ability to perform age-related activities of daily living and/or impairing functional abilities;]

9. Chelation therapy. The administration of FDA-approved chelating agents is covered for any of the following conditions:

[A. Genetic or hereditary hemochromatosis;
B. Lead overload in cases of acute or long-term lead exposure;
C. Secondary hemochromatosis due to chronic iron overload due to transfusion-dependent anemias (e.g., Thalassemias, Cooley’s anemia, sickle cell anemia, sideroblastic anemia);
D. Copper overload in patients with Wilson’s disease;
E. Arsenic, mercury, iron, copper, or gold poisoning when long-term exposure to and toxicity has been confirmed through lab results or clinical findings consistent with metal toxicity;
F. Aluminum overload in chronic hemodialysis patients;
G. Emergency treatment of hypercalcemia;
H. Prophylaxis against doxorubicin-induced cardiomyopathy;
I. Internal plutonium, americium, or curium contamination; or
J. Cystinuria;]

10. Chiropractic services. Chiropractic—manipulation and adjunct therapeutic procedures/modalities (e.g., mobilization, therapeutic exercise, traction) are covered when all of the following conditions are met:

[A. A neuromusculoskeletal condition is diagnosed that may be relieved by standard chiropractic treatment in order to restore optimal function;
B. Chiropractic care is being performed by a licensed doctor of chiropractic who is practicing within the scope of his/her license as defined by state law;
C. The individual is involved in a treatment program that clearly documents all of the following:
(I) A prescribed treatment program that is expected to result in significant therapeutic improvement over a clearly defined period of time;
(II) The symptoms being treated;
(III) Diagnostic procedures and results;
(IV) Frequency, duration, and results of planned treatment modalities;
(V) Anticipated length of treatment plan with identification of quantifiable, attainable short-term and long-term goals; and
(VI) Demonstrated progress toward significant functional gains and/or improved activity tolerances;
D. Following previous successful treatment with chiropractic care, acute exacerbation or re-injury are covered when all of the following criteria are met:
(II) The member reached maximal therapeutic benefit with prior chiropractic treatment;
(II) The member was compliant with a self-directed homecare program;
(III) Significant therapeutic improvement is expected with continued treatment; and
(IV) The anticipated length of treatment is expected to be short-term (e.g., no more than six (6) visits within a three- (3-) week period;)

11. Clinical trials. Routine member care costs incurred as the result of a Phase I, II, III, or IV clinical trial that is conducted in relation to the prevention, detection, or treatment of cancer or other life-threatening disease or condition are covered when—

A. The study or investigation is conducted under an investigational new drug application reviewed by the FDA; or
B. Is a drug trial that is exempt from having such an investigational new drug application. Life-threatening condition means any disease or condition from which the likelihood of death is probable unless the course of the disease or condition is interrupted; and
C. Routine patient care costs include all items and services consistent with the coverage provided in plan benefits that would otherwise be covered for a member not enrolled in a clinical trial. Routine patient care costs do not include the investigational item, device, or service itself; items and services that are provided solely to satisfy data collection and analysis needs and are not used in the direct clinical management of the member; or a service that is clearly inconsistent with widely accepted and established standards of care for a particular diagnosis;
D. The member must be eligible to participate in the clinical trial according to the trial protocol with respect to treatment of cancer or other life-threatening disease or condition; and
E. The clinical trial must be approved or funded by one (1) of the following:

(I) National Institutes of Health (NIH);
(II) Centers for Disease Control and Prevention (CDC);
(III) Agency for Health Care Research and Quality;
(IV) Centers for Medicare & Medicaid Services (CMS);
(V) A cooperative group or center of any of the previously named agencies or the Department of Defense or the Department of Veterans Affairs;
(VI) A qualified non-governmental research entity identified in the guidelines issued by the National Institutes of Health for center support grants; or
(VII) A study or investigation that is conducted by the Department of Veterans Affairs, the Department of Defense, or the Department of Energy and has been reviewed and approved to be comparable to the system of peer review of studies and investigations used by the NIH and assures unbiased review of the highest scientific standards by qualified individuals who have no interest in the outcome of the review;

12. Cochlear implant. Unilateral (monaural) or bilateral (bilateral) cochlear implantation and necessary replacement batteries are covered for a member with bilateral, pre- or post-linguistic, sensorineural, moderate-to-profound hearing impairment when there is reasonable expectation that a significant benefit will be achieved from the device and when the following age-specific criteria are met:

[aUDitory brainstem implant]

[A. Auditory brainstem implant. Auditory brainstem implant (ABI) covered for the diagnosis of neurofibromatosis type II, von Recklinghausen’s disease, or when a member is undergoing bilateral removal of tumors of the auditory nerves, and it is anticipated that the member will become completely deaf as a result of the surgery, or the member had bilateral auditory nerve tumors removed and is now bilaterally deaf;
(II) For an adult (age eighteen (18) years or older) with BOTH of the following:
(a) Bilateral, severe to profound sensorineural hearing loss determined by a pure-tone average of seventy (70) decibels (dB) hearing loss or greater at five hundred (500) hertz (Hz), one thousand (1000) Hz, and two thousand (2000) Hz; and

(b) Member has limited benefit from appropriately fitted binaural hearing aids. Limited benefit from amplification is defined by test scores of forty percent (40%) correct or less in best-aided listening condition on open-set sentence cognition (e.g., Central Institute for the Deaf (CID) sentences, Hearing in Noise Test (HINT) sentences, and Consonant-Nucleus-Consonant (CNC) test); and

(ii) For a child age twelve (12) months to seventeen (17) years, eleven (11) months with both of the following:

(a) Profound, bilateral sensorineural hearing loss with thresholds of ninety (90) dB or greater at one thousand (1000) Hz; and

(b) Limited or no benefit from a three- (3-) month trial of appropriately fitted binaural hearing aids;

(iii) For children four (4) years of age or younger, with one (1) of the following:

(a) Failure to reach developmentally appropriate auditory milestones measured using the Infant-Toddler Meaningful Auditory Integration Scale, the Meaningful Auditory Integration Scale, or the Early Speech Perception test; or

(b) Less than twenty percent (20%) correct on open-set word recognition test Multisyllabic Lexical Neighborhood Test (MLNT) in conjunction with appropriate amplification and participation in intensive aural habilitation over a three- (3-) to six- (6-) month period;

(iv) For children older than four (4) years of age with one (1) of the following:

(a) Less than twelve percent (12%) correct on the Phonetically Balanced-Kindergarten Test; or

(b) Less than thirty percent (30%) correct on the HINT for children, the open-set Multisyllabic Lexical Neighborhood Test (MLNT) or Lexical Neighborhood Test (LNT), depending on the child’s cognitive ability and linguistic skills; and

(v) A three- (3-) to six- (6-) month hearing aid trial has been undertaken by a child without previous experience with hearing aids;

B. Radiologic evidence of cochlear ossification;

C. The following additional medical necessity criteria must also be met for uniaural (monaural) or binaural (bilateral) cochlear implantation in adults and children:

(I) Member must be enrolled in an educational program that supports listening and speaking with aided hearing;

(II) Member must have had an assessment by an audiologist and from an otolaryngologist experienced in this procedure indicating the likelihood of success with this device;

(III) Member must have no medical contraindications to cochlear implantation (e.g., cochlear aplasia, active middle ear infection); and

(IV) Member must have arrangements for appropriate follow-up care, including the speech therapy required to take full advantage of this device;

D. A second cochlear implant is covered in the contralateral (opposite) ear as medically necessary in an individual with an existing unilateral cochlear implant when the hearing aid in the contralateral ear produces limited or no benefit;

E. The replacement of an existing cochlear implant is covered when either of the following criteria is met:

(I) Currently used component is no longer functional and cannot be repaired; or

(II) Currently used component renders the implant recipient unable to adequately and/or safely perform his/her age-appropriate activities of daily living; and

F. Post-cochlear or ABI rehabilitation program (aural rehabilitation) is covered to achieve benefit from a covered device;]


A. Dental care is covered for the following:

(I) Treatment to reduce trauma and restorative services limited to dental implants only when the result of accidental injury to sound natural teeth and tissue that are viable, functional, and free of disease. Treatment must be initiated within sixty (60) days of accident; and

(II) Restorative services limited to dental implants when needed as a result of [cancerous or non-cancerous] tumors and cysts, cancer, and post-surgical sequelae.

B. The administration of general anesthesia, monitored anesthesia care, and hospital charges for dental care are covered for children younger than five (5) years, the severely disabled, or a person with a medical or behavioral condition that requires hospitalization when provided in a network or non-network hospital or surgical center;

14. Diabetes Self-Management Education;

15. Dialysis is covered when received through a network provider;

16. Durable medical equipment (DME) is covered when ordered by a provider to treat an injury or illness. DME includes, but is not limited to, the following:

A. Insulin pumps;

B. Oxygen;

C. Augmentative communication devices;

D. Manual and powered mobility devices;

E. Disposable supplies that do not withstand prolonged use and are periodically replaced, including, but not limited to, the following:

(I) Colostomy and ureterostomy bags;

(II) Prescription compression stockings limited to two (2) pairs or four (4) individual stockings per plan year;

F. Blood pressure cuffs/monitors with a diagnosis of diabetes;

G. Repair and replacement of DME is covered when any of the following criteria are met:

(I) Repairs, including the replacement of essential accessories, which are necessary to make the item or device serviceable;

(II) Routine wear and tear of the equipment renders it nonfunctional and the member still requires the equipment; or

(III) The provider has documented that the condition of the member changes or if growth-related;

17. Emergency room services. Coverage is for emergency medical conditions. If a member is admitted to the hospital, s/he may be required to transfer to network facility for maximum benefit.

Hospital and ancillary charges are paid as a network benefit;

18. Eye glasses and contact lenses. Coverage limited to charges incurred in connection with the fitting of eye glasses or contact lenses for initial placement within one (1) year following cataract surgery;

19. Foot care (trimming of nails, corns, or calluses). Foot care services are covered when administered by a provider and—

A. When associated with systemic conditions that are significant enough to result in severe circulatory insufficiency or areas of desensitization in the lower extremities including, but not limited to, any of the following:

(I) Diabetes mellitus;

(II) Peripheral vascular disease; or

(III) Peripheral neuropathy.

(IV) Evaluation/debridement of mycotic nails, in the absence of a systemic condition, when both of the following conditions are met:
(a) Pain or secondary infection resulting from the thickening and dystrophy of the infected toenail plate; and
(b) If the member is ambulatory, pain markedly limits ambulation;

20. Genetic counseling. Pre-test and post-test genetic counseling with a provider or a licensed or certified genetic counselor are covered when a member is recommended for covered heritable genetic testing.

[A. Genetic counseling in connection with pregnancy management is covered only for evaluation of any of the following:
   (I) Couples who are closely related genetically (e.g., consanguinity, incest);
   (II) Familial cancer disorders;
   (III) Individuals recognized to be at increased risk for genetic disorders;
   (IV) Infertility cases where either parent is known to have a chromosomal abnormality;
   (V) Primary amenorrhea, azoopermia, abnormal sexual development, or failure in developing secondary sexual characteristics;
   (VI) Mother is a known, or presumed carrier of an X-linked recessive disorder;
   (VII) One (1) or both parents are known carriers of an autosomal recessive disorder;
   (VIII) Parents of a child born with a genetic disorder, birth defect, inborn error of metabolism, or chromosome abnormality;
   (IX) Parents of a child with intellectual developmental disorders, autism, developmental delays, or learning disabilities;
   (X) Pregnant women who, based on prenatal ultrasound tests or an abnormal multiple marker screening test, maternal serum alpha-fetoprotein (AFP) test, test for sickle cell anemia, or tests for other genetic abnormalities have been told their pregnancy may be at increased risk for complications or birth defects;
   (XI) Pregnant women age thirty-five (35) years or older at delivery;
   (XII) Pregnant women, or women planning pregnancy, exposed to potentially teratogenic, mutagenic, or carcinogenic agents such as chemicals, drugs, infections, or radiation;
   (XIII) Previous unexplained stillbirth or repeated (three (3) or more; two (2) or more among infertile couples) first-trimester miscarriages, where there is suspicion of parental or fetal chromosome abnormalities; or
   (XIV) When contemplating pregnancy, either parent affected with an autosomal dominant disorder;]

21. Genetic testing.

A. Genetic testing is covered to establish a molecular diagnosis of an inheritable disease when all of the following criteria are met:
   (I) The member displays clinical features or is at direct risk of inheriting the mutation in question (pre-symptomatic);
   (II) The result of the test will directly impact the treatment being delivered to the member;
   (III) The testing method is considered scientifically valid for identification of a genetically-linked heritable disease; and
   (IV) After history, physical examination, pedigree analysis, genetic counseling, and completion of conventional diagnostic studies, a definitive diagnosis remains uncertain.

B. Genetic testing for the breast cancer susceptibility gene (BRCA) when family history is present;

22. Hair analysis. Chemical hair analysis is covered for the diagnosis of suspected chronic arsenic poisoning. Other purposes are considered experimental and investigational;

23. Hair prostheses. Prostheses and expenses for scalp hair prostheses worn for hair loss are covered for alopecia areata or alopecia totalis for children eighteen (18) years of age or younger. The annual maximum is two hundred dollars ($200), and the lifetime maximum is three thousand two hundred dollars ($3,200);

24. Hearing aids (per ear). Hearing aids are covered once every two (2) years for conductive hearing loss unresponsive to medical or surgical interventions, sensorineural hearing loss, and mixed hearing loss.

[A. Prior to receiving a hearing aid members must receive—
   (I) A medical exam by a physician or other qualified provider to identify any medically treatable conditions that may affect hearing; and
   (II) A comprehensive hearing test to assess the need for hearing aids conducted by a certified audiologist, hearing instrument specialist, or other provider licensed or certified to administer this test.

B. Covered once every two (2) years.] If the cost of one (1) hearing aid exceeds the amount listed below, member is also responsible for charges over that amount.

[(III)]
(A. Conventional: one thousand dollars ($1,000).
(B. Programmable: two thousand dollars ($2,000).
(C. Digital: two thousand five hundred dollars ($2,500).
(D. Bone Anchoring Hearing Aid (BAHA): three thousand five hundred dollars ($3,500);

25. Hearing testing. One (1) hearing test per year. Additional hearing tests are covered if recommended by provider;

26. Home health care. Skilled home health nursing care is covered for members who are homebound because of injury or illness (i.e., the member leaves home only with considerable and taxing effort, and absences from home are infrequent or of short duration, or to receive medical care). Services must be performed by a registered nurse or licensed practical nurse, licensed therapist, or a registered dietitian. Covered services include:

A. Home visits instead of visits to the provider’s office that do not exceed the usual and customary charge to perform the same service in a provider’s office;
B. Intermittent nurse services. Benefits are paid for only one (1) nurse at any one (1) time, not to exceed four (4) hours per twenty-four (24-) hour period;
C. Nutrition counseling provided by or under the supervision of a registered dietitian;
D. Physical, occupational, respiratory, and speech therapy provided by or under the supervision of a licensed therapist;
E. Medical supplies, drugs, or medication prescribed by provider, and laboratory services to the extent that the plan would have covered them under this plan if the covered person had been in a hospital;
F. A home health care visit is defined as—
   (I) A visit by a nurse providing intermittent nurse services (each visit includes up to a four-hour (4-) hour consecutive visit in a twenty-four (24-) hour period if clinical eligibility for coverage is met) or a single visit by a therapist or a registered dietitian; and

G. Benefits cannot be provided for any of the following:
   (I) Homemaker or housekeeping services;
   (II) Supportive environment materials such as handrails, ramps, air conditioners, and telephones;
   (III) Services performed by family members or volunteer workers;
   (IV) “Meals on Wheels” or similar food service;
   (V) Separate charges for records, reports, or transportation;
   (VI) Expenses for the normal necessities of living such as food, clothing, and household supplies; and
   (VII) Legal and financial counseling services, unless otherwise covered under this plan;
27. Hospice care and palliative services (inpatient or outpatient). Includes bereavement and respite care. Hospice care services, including pre-hospice evaluation or consultation, are covered when the individual is terminally ill (and expected to live six (6) months or less, potentially curative treatment for the terminal illness is not part of the prescribed plan of care, the individual or appointed designee has formally consented to hospice care (i.e., care directed mostly toward palliative care and symptom management), and the hospice services are provided by a certified/accredited hospice agency with care available twenty-four (24) hours per day, seven (7) days per week; 

[A. When the above criteria are met, the following hospice care services are covered:

(I) Assessment of the medical and social needs of the terminally ill person, and a description of the care to meet those needs;

(II) Inpatient care in a facility when needed for pain control and other acute and chronic symptom management, psychological and dietary counseling, physical or occupational therapy, and part-time home health care services;

(III) Outpatient care for other services as related to the terminal illness, which include services of a physician, physical or occupational therapy, and nutrition counseling provided by or under the supervision of a registered dietitian; and

(IV) Bereavement counseling benefits which are received by a member’s close relative when directly connected to the member’s death and bundled with other hospice charges. The services must be furnished within twelve (12) months of death.]

28. Hospital (includes inpatient, outpatient, and surgical centers).

A. The following benefits are covered:

(I) Semi-private room and board. For network charges, this rate is based on network repricing. For non-network charges, any charge over a semi-private room charge will be a covered expense only when clinical eligibility for coverage is met. If the hospital has no semi-private rooms, the plan will allow the private room rate subject to usual, customary, and reasonable charges or the network rate, whichever is applicable;

(II) Intensive care unit room and board;

(III) Surgery, therapies, and ancillary services including, but not limited to:

(a) Cornea transplant;

(b) Coverage for breast reconstruction surgery or prostheses following mastectomy and lumpectomy is available to both females and males. A diagnosis of breast cancer is not required for breast reconstruction services to be covered, and the timing of reconstructive services is not a factor in coverage;

(c) Sterilization for the purpose of birth control is covered;

(d) Cosmetic/reconstructive surgery is covered to repair a functional disorder caused by disease or injury;

(e) Cosmetic/reconstructive surgery is covered to repair a congenital defect or abnormality for a member younger than nineteen (19) years; and

(f) Blood, blood plasma, and plasma expanders are covered, when not available without charge;

(IV) Inpatient mental health services [are covered when authorized by a physician for treatment of a mental health disorder. Inpatient mental health services are covered, subject to all of the following; and]

[a] Member must be ill in more than one (1) area of daily living to such an extent that s/he is rendered dysfunctional and requires the intensity of an inpatient setting for treatment. Without such inpatient treatment, the member’s condition would deteriorate;

[b] The member’s mental health disorder must be treatable in an inpatient facility;

(c) The member’s mental health disorder must meet diagnostic criteria as described in the most recent edition of the American Psychiatric Association Diagnostic and Statistical Manual (DSM). If outside of the United States, the member’s mental health disorder must meet diagnostic criteria established and commonly recognized by the medical community in that region;

[d] The attending provider must be a psychiatrist. If the admitting provider is not a psychiatrist, a psychiatrist must be attending to the member within twenty-four (24) hours of admittance. Such psychiatrist must be United States board-eligible or board-certified. If outside of the United States, inpatient services must be provided by an individual who has received a diploma from a medical school recognized by the government agency in the country where the medical school is located. The attending provider must meet the requirements, if any, set out by the foreign government or regionally-recognized licensing body for treatment of mental health disorders;

(e) Day treatment (partial hospitalization) for mental health services means a day treatment program that offers intensive, multidisciplinary services provided on less than a full-time basis. The program is designed to treat patients with serious mental or nervous disorders and offers major diagnostic, psychosocial, and prevocational modalities. Such programs must be a less-restrictive alternative to inpatient treatment; and

(f) Mental health services received in a residential treatment facility that is licensed by the state in which it operates and provides treatment for mental health disorders is covered. This does not include services provided at a group home. If outside of the United States, the residential treatment facility must be licensed or approved by the foreign government or an accreditation or licensing body working in that foreign country; and

(V) Outpatient mental health services [are covered if the member is at a therapeutic medical or mental health facility and treatment includes measurable goals and continued progress toward functional behavior and termination of treatment. Continued coverage may be denied when positive response to treatment is not evident. Treatment must be provided by one (1) of the following;]

[a] A United States board-eligible or board-certified psychiatrist licensed in the state where the treatment is provided;

(b) A therapist with a doctorate or master’s degree that denotes a specialty in psychiatry (Psy.D.);

(c) A state-licensed psychologist;

(d) A state-licensed or certified social worker practicing within the scope of his or her license or certification; or

(e) Licensed professional counselor;]

29. Infusions are covered when received through a network provider. Medications (specialty and non-specialty) that can be safely obtained through a pharmacy and which may be self-administered are not a medical plan benefit but are covered as part of the pharmacy benefit;

30. Injections. See preventive services for coverage of vaccinations. See contraception and sterilization for coverage of birth control injections. Medications (specialty and non-specialty) that can be safely obtained through a pharmacy and which may be self-administered are not a medical plan benefit but are covered as part of the pharmacy benefit;
31. Lab, X-ray, and other diagnostic procedures. Outpatient diagnostic services are covered when tests or procedures are performed for a specific symptom and to detect or monitor a condition. Professional charges for automated lab services performed by an out-of-network provider are not covered.

32. Maternity coverage. Prenatal and postnatal care is covered. Routine prenatal office visits and screenings recommended by the Health Resources and Services Administration are covered at one hundred percent (100%). Other care is subject to applicable deductibles and coinsurance. Newborns and their mothers are allowed hospital stays of at least forty-eight (48) hours after vaginal birth and ninety-six (96) hours after cesarean section birth. If discharge occurs earlier than specific time periods, the plan shall provide coverage for post discharge care that shall consist of a two- (2-) visit minimum, at least one (1) in the home.

33. Nutritional counseling. Individualized nutritional evaluation and counseling for the management of any medical condition for which appropriate diet and eating habits are essential to the overall treatment program is covered when ordered by a physician or physician extender and provided by a licensed health-care professional (e.g., a registered dietitian).

34. Nutrition therapy.

A. Nutrition therapy is covered only when the following criteria are met:

(I) Nutrition therapy is the sole source of nutrients or a significant percentage of the daily caloric intake;

(II) Nutrition therapy is used in the treatment of, or in association with, a demonstrable disease, condition, or disorder;

(III) Nutrition therapy is necessary to sustain life or health;

(IV) Nutrition therapy is prescribed by a provider;

and

(V) Nutrition therapy is managed, monitored, and evaluated on an on-going basis, by a provider.

B. Only the following types of nutrition therapy are covered:

(I) Enteral Nutrition (EN). EN is the provision of nutritional requirements via the gastrointestinal tract. EN can be taken orally or through a tube into the stomach or small intestine;

(II) Parenteral Nutrition Therapy (PN) and Total Parenteral Nutrition (TPN). PN is liquid nutrition administered through a vein to provide part of daily nutritional requirements. TPN is a type of PN that provides all daily nutrient needs. PN or TPN are covered when the member’s nutritional status cannot be adequately maintained on oral or enteral feedings.

(III) Intradialytic Parenteral Nutrition (IDPN). IDPN is a type of PN that is administered to members on chronic hemodialysis during dialysis sessions to provide most nutrient needs. IDPN is covered when the member is on chronic hemodialysis and nutritional status cannot be adequately maintained on oral or enteral feedings.

35. Office visit. Member encounter with a provider for health care, mental health, or substance use disorder in an office, clinic, or ambulatory care facility is covered based on the service, procedure, or related treatment plan.

36. Oral surgery is covered for injury, tumors, or cysts. Oral surgery includes, but is not limited to, reduction of fractures and dislocation of the jaws; external incision and drainage of cellulites; incision of accessory sinuses, salivary glands, or ducts; excision of exostosis of jaws and hard palate; and frenectomy. Treatment must be initiated within sixty (60) days of accident. No coverage for dental care, including oral surgery, as a result of poor dental hygiene. Extractions of bony or partial bony impactions are excluded.

37. Orthognathic or Jaw Surgery. Orthognathic or jaw surgery is covered when one (1) of the following conditions is documented and diagnosed:

A. Acute traumatic injury, and post-surgical sequela;

B. Cancerous or non-cancerous tumors, cysts, cancer, and post-surgical sequela;

C. Cleft lip/palate (for cleft lip/palate related jaw surgery); or

D. Physical or physiological abnormality [when one (1) of the following criteria is met:]

(I) Anteroposterior Discrepancies—

(a) Maxillary/Mandibular incisor relationship: over jet of 5mm or more, or a 0 to a negative value (norm 2mm);

(b) Maxillary/Mandibular anteroposterior molar relationship discrepancy of 4mm or more (norm 0 to 1mm); or

(c) These values represent two (2) or more standard deviation from published norms;

(II) Vertical Discrepancies—

(a) Presence of a vertical facial skeletal deformity which is two (2) or more standard deviations from published norms for accepted skeletal landmarks;

(b) Open bite with no vertical overlap of anterior teeth or unilateral or bilateral posterior open bite greater than 2mm;

(c) Deep overbite with impingement or irritation of buccal or lingual soft tissues of the opposing arch; or

(d) Supraeruption of a dentoalveolar segment due to lack of occlusion;

(III) Transverse Discrepancies—

(a) Presence of a transverse skeletal discrepancy which is two (2) or more standard deviations from published norms; or

(b) Total bilateral maxillary palatal cusp to mandibular fossa discrepancy of 4mm or greater, or a unilateral discrepancy of 3mm or greater, given normal axial inclination of the posterior teeth; or

(IV) Asymmetries—

(a) Anteroposterior, transverse, or lateral asymmetries greater than 3mm with concomitant occlusal asymmetry;

(V) Masticatory (chewing) and swallowing dysfunction due to malocclusion (e.g., inability to incise or chew solid foods, choking on incompletely masticated solid foods, damage to soft tissue during mastication, malnutrition); and

(VI) Speech impairment; or

(VII) Obstructive sleep apnea or airway dysfunction;]
38. Orthotics.
   A. Ankle–Foot Orthosis (AFO) and Knee–Ankle–Foot Orthosis (KAFO).
      (I) Basic coverage criteria for AFO and KAFO used during ambulation are as follows:
      (a) AFO is covered when used in ambulation for members with weakness or deformity of the foot and ankle, which require stabilization for medical reasons, and have the potential to benefit functionally;
      (b) KAFO is covered when used in ambulation for members when the following criteria are met:
         I. Member is covered for AFO; and
         II. Additional knee stability is required; and
      (c) AFO and KAFO that are molded-to-patient-model, or custom-fabricated, are covered when used in ambulation, only when the basic coverage criteria and one (1) of the following criteria are met:
         I. The member could not be fitted with a prefabricated AFO;
         II. AFO or KAFO is expected to be permanent or for more than six (6) months duration;
         III. Knee, ankle, or foot must be controlled in more than one (1) plane;
         IV. There is documented neurological, circulatory, or orthopedic status that requires custom fabricating over a model to prevent tissue injury; or
         V. The member has a healing fracture which lacks normal anatomical integrity or anthropometric proportions.
      (II) AFO and KAFO Not Used During Ambulation.
      (a) AFO and KAFO not used in ambulation are covered if the following criteria are met:
         I. Passive range of motion test was measured with agonimeter and documented in the medical record;
         II. Documentation of an appropriate stretching program administered under the care of provider or caregiver;
         III. Plantar flexion contracture of the ankle with dorsiflexion on passive range of motion testing of at least ten degrees (10°) (i.e., a non-fixed contracture);
         IV. Reasonable expectation of the ability to correct the contracture;
         V. Contracture is interfering or expected to interfere significantly with the patient’s functional abilities; and
         VI. Used as a component of a therapy program which includes active stretching of the involved muscles and/or tendons; or
         VII. Member has plantar fasciitis.
      (b) Replacement interface for AFO or KAFO is covered only if member continues to meet coverage criteria and is limited to a maximum of one (1) per six (6) months.
   B. Cast Boot, Post-Operative Sandal or Shoe, or Healing Shoe. A cast boot, post-operative sandal or shoe, or healing shoe is covered for one (1) of the following indications:
      (I) To protect a cast from damage during weight-bearing activities following injury or surgery;
      (II) To provide appropriate support and/or weight-bearing surface to a foot following surgery;
      (III) To promote good wound care and/or healing via appropriate weight distribution and foot protection; or
      (IV) When the patient is currently receiving treatment for lymphedema and the foot cannot be fitted into conventional footwear.
   C. Cranial Orthoses. Cranial orthosis is covered for Synostotic and Non-Synostotic Plagiocephaly. Plagiocephaly is an asymmetrically shaped head. Synostotic Plagiocephaly is due to premature closure of cranial sutures. Non-Synostotic Plagiocephaly is from positioning or deformation of the head. Cranial orthosis is the use of a special helmet or band on the head which aids in molding the shape of the cranium to normal. Initial reimbursement shall cover any subsequent revisions.
   D. Elastic Supports. Elastic supports are covered when prescribed for one (1) of the following indications:
      (I) Severe or incapacitating vascular problems, such as acute thrombophlebitis, massive venous stasis, or pulmonary embolism;
      (II) Venous insufficiency;
      (III) Varicose veins;
      (IV) Edema of lower extremities;
      (V) Edema during pregnancy; or
      (VI) Lymphedema.
   E. Footwear Incorporated Into a Brace for Members with Skeletally Mature Feet. Footwear incorporated into a brace must be billed by the same supplier billing for the brace. The following types of footwear incorporated into a brace are covered:
      (I) Orthopedic footwear;
      (II) Other footwear such as high top, depth inlay, or custom;
      (III) Heel replacements, sole replacements, and shoe transfers involving shoes on a brace;
      (IV) Inserts for a shoe that is an integral part of a brace and are required for the proper functioning of the brace; or
      (V) Other shoe modifications if they are on a shoe that is an integral part of a brace and are required for the proper functioning of the brace.
   F. Foot Orthoses. Custom, removable foot orthoses are covered for members who meet the following criteria:
      (I) Member with skeletally mature feet who has any of the following conditions:
         (a) Acute plantar fasciitis;
         (b) Acute sport-related injuries with diagnoses related to inflammatory problems such as bursitis or tendinitis;
         (c) Calcaneal bursitis (acute or chronic);
         (d) Calcaneal spurs (heel spur);
         (e) Conditions related to diabetes;
         (f) Inflammatory conditions (e.g., sesamoiditis, submetatarsal bursitis, synovitis, tenosynovitis, synovial cyst, osteomyelitis, and plantar fascial fibromatosis);
         (g) Medial osteoarthritis of the knee;
         (h) Musculoskeletal/arthropathic deformities including deformities of the joint or skeleton that impairs walking in a normal shoe (e.g., bunions, hallux valgus, talipes deformities, pes deformities, or anomalies of toes);
         (i) Neurologically impaired feet including neuroma, tarsal tunnel syndrome, ganglionic cyst;
         (j) Neuropathies involving the feet, including those associated with peripheral vascular disease, diabetes, carcinoma, drugs, toxins, and chronic renal disease; or
         (k) Vascular conditions including ulceration, poor circulation, peripheral vascular disease, Buerger’s disease (thromboangiitis obliterans), and chronic thrombophlebitis;
      (II) Member with skeletally immature feet who has any of the following conditions:
         (a) Hallux valgus deformities;
         (b) In-toe or out-toe gait;
         (c) Musculoskeletal weakness such as pronation or pes planus;
         (d) Structural deformities such as tarsal coalition;
         (e) Torsional conditions such as metatarsus adductus, tibial torsion, or femoral torsion.
   G. Helmets. Helmets are covered when cranial protection is required due to a documented medical condition that makes the member susceptible to injury during activities of daily living.
   H. Hip Orthosis. Hip orthosis is covered for one (1) of the following indications:
      (I) To reduce pain by restricting mobility of the hip;
      (II) To facilitate healing following an injury to the hip or
related soft tissues;
(III) To facilitate healing following a surgical procedure of the hip or related soft tissue; or
(IV) To otherwise support weak hip muscles or a hip deformity.
I. Knee Orthosis. Knee orthosis is covered for one (1) of the following indications:
(I) To reduce pain by restricting mobility of the knee;
(II) To facilitate healing following an injury to the knee or related soft tissues;
(III) To facilitate healing following a surgical procedure on the knee or related soft tissue; or
(IV) To otherwise support weak knee muscles or a knee deformity.
J. Orthopedic Footwear for Diabetic Members.
(I) Orthopedic footwear, therapeutic shoes, inserts, or modifications to therapeutic shoes are covered for diabetic members if any following criteria are met:
(a) Previous amputation of the other foot or part of either foot;
(b) History of previous foot ulceration of either foot;
(c) History of pre-ulcerative calluses of either foot;
(d) Peripheral neuropathy with evidence of callus formation of either foot;
(e) Foot deformity of either foot; or
(f) Poor circulation in either foot.
(II) Coverage is limited to one (1) of the following within one (1) year:
(a) One (1) pair of custom molded shoes (which includes inserts provided with these shoes) and two (2) additional pairs of inserts;
(b) One (1) pair of depth shoes and three (3) pairs of inserts (not including the non-customized removable inserts provided with such shoes); or
(c) Up to three (3) pairs of inserts not dispensed with diabetic shoes if the supplier of the shoes verifies in writing that the patient has appropriate footwear into which the insert can be placed.
K. Orthotic-Related Supplies. Orthotic-related supplies are covered when necessary for the function of the covered orthotic device.
L. Spinal Orthoses. A thoracic-lumbar-sacral orthosis, lumbar orthosis, lumbar-sacral orthosis, and cervical orthosis are covered for the following indications:
(I) To reduce pain by restricting mobility of the trunk;
(II) To facilitate healing following an injury to the spine or related soft tissues;
(III) To facilitate healing following a surgical procedure of the spine or related soft tissue; or
(IV) To otherwise support weak spinal muscles or a deformed spine.
M. Trusses. Trusses are covered when a hernia is reducible with the application of a truss.
N. Upper Limb Orthosis. Upper limb orthosis is covered for the following indications:
(I) To reduce pain by restricting mobility of the joint(s);
(II) To facilitate healing following an injury to the joint(s) or related soft tissues; or
(III) To facilitate healing following a surgical procedure of the joint(s) or related soft tissue.
O. Orthotic Device Replacement. When repairing an item that is no longer cost-effective and is out of warranty, the plan will consider replacing the item subject to review of medical necessity and life expectancy of the device.
39. Preventive services.
A. Services recommended by the U.S. Preventive Services Task Force (categories A and B).
B. Vaccinations recommended by the Advisory Committee on Immunization Practices of the Centers for Disease Control and Prevention.
C. Preventive care and screenings for infants, children, and adolescents supported by the Health Resources and Services Administration.
D. Preventive care and screenings for women supported by the Health Resources and Services Administration.
E. Preventive exams and other services ordered as part of the exam. For benefits to be covered as preventive, they must be coded by the provider as routine, without indication of an injury or illness.
F. Cancer screenings. One (1) per calendar year. Additional screenings beyond one (1) per calendar year covered as diagnostic unless otherwise specified—
(I) Mammograms—no age limit. Standard two-dimensional (2D) breast mammography and breast tomosynthesis (three-dimensional (3D) mammography);
(II) Pap smears—no age limit;
(III) Prostate—no age limit; and
(IV) Colorectal screening—no age limit.
G. Online weight management program offered through the plan’s exclusive provider arrangement;
40. Prostheses (prosthetic devices). Basic equipment that meets medical needs. Repair and replacement is covered due to normal wear and tear, if there is a change in medical condition, or if growth-related;
41. Pulmonary rehabilitation. Comprehensive, individualized, goal-directed outpatient pulmonary rehabilitation covered for pre- and post-operative intervention for lung transplantation and lung volume reduction surgery (LVRS) or when all of the following apply:
A. Member has a reduction of exercise tolerance that restricts the ability to perform activities of daily living (ADL) or work;
B. Member has chronic pulmonary disease (including asthma, emphysema, chronic bronchitis, chronic airflow obstruction, cystic fibrosis, alpha-1 antitrypsin deficiency, pneumonia, asbestosis, radiation pneumonitis, pulmonary fibrosis, pulmonary alveolar proteinosis, pulmonary hemosiderosis, fibrosing alveolitis), or other conditions that affect pulmonary function such as ankylosing spondylitis, scoliosis, myasthenia gravis, muscular dystrophy, Guillain-Barré syndrome, or other infective polyneuritis, sarcoidosis, paralysis of diaphragm, or bronchopulmonary dysplasia; and
C. Member has a moderate to moderately severe functional pulmonary disability, as evidenced by either of the following, and does not have any concomitant medical condition that would otherwise imminently contribute to deterioration of pulmonary status or undermine the expected benefits of the program (e.g., symptomatic coronary artery disease, congestive heart failure, myocardial infarction within the last six (6) months, dysrhythmia, active joint disease, claudication, malignancy):
(I) A maximal pulmonary exercise stress test under optimal bronchodilatory treatment which demonstrates a respiratory limitation to exercise with a maximal oxygen uptake (VO2 max) equal to or less than twenty milliliters per kilogram per minute (20 mL/kg/min), or about five (5) metabolic equivalents (METs); or
(II) Pulmonary function tests showing that either the Forced Expiratory Volume in One Second (FEV1), Forced Vital Capacity (FVC), FEV1/FVC, or Diffusing Capacity of the Lung for Carbon Monoxide (DLCO) is less than sixty percent (60%) of that predicted;
42. Skilled Nursing Facility. Skilled nursing facility services are covered up to one hundred twenty (120) days per calendar year;
43. Telehealth Services. Telehealth services are covered for the diagnosis, consultation, or treatment of a member on the same basis that the service would be covered when it is delivered in person;
44. Therapy. Physical, occupational, and speech therapy are covered when prescribed by a provider and subject to the provisions below:
A. Physical therapy.
(I) Physical therapy must meet the following criteria:
(a) The program is designed to improve lost or impaired physical function or reduce pain resulting from illness, injury, congenital defect, or surgery;
(b) The program is expected to result in significant therapeutic improvement over a clearly defined period of time; and
(c) The program is individualized, and there is documentation outlining quantifiable, attainable treatment goals;

B. Occupational therapy must meet the following criteria:
(I) The program is designed to improve or compensate for lost or impaired physical functions, particularly those affecting activities of daily living, resulting from illness, injury, congenital defect, or surgery;
(II) The program is expected to result in significant therapeutic improvement over a clearly defined period of time; and
(III) The program is individualized, and there is documentation outlining quantifiable, attainable treatment goals;

C. Speech therapy.
(I) All of the following criteria must be met for coverage of speech therapy:
(a) The therapy requires one-to-one intervention and supervision of a speech-language pathologist;
(b) The therapy plan includes specific tests and measures that will be used to document significant progress every two (2) weeks;
(c) Meaningful improvement is expected;
(d) The therapy includes a transition from one-to-one supervision to a self- or caregiver- provided maintenance program upon discharge; and
(e) One (1) of the following:
   I. Member has severe impairment of speech-language; and an evaluation has been completed by a certified speech-language pathologist that includes age-appropriate standardized tests to measure the extent of the impairment, performance deviation, and language and pragmatic skill assessment levels; or
   II. Member has a significant voice disorder that is the result of anatomic abnormality, neurological condition, or injury (e.g., vocal nodules or polyps, vocal cord paresis or paralysis, post-operative vocal cord surgery);

45. Transplants. Stem cell, kidney, liver, heart, lung, pancreas, small bowel, or any combination are covered. Includes services related to organ procurement and donor expenses if not covered under another plan. Member must contact medical plan for arrangements.

A. Network includes travel and lodging allowance for the transplant recipient and an immediate family travel companion when the transplant facility is more than fifty (50) miles from the recipient’s residence. If the recipient is younger than age nineteen (19) years, travel and lodging is covered for both parents. The transplant recipient must be with the travel companion or parent(s) travel expense to be reimbursable. Combined travel and lodging expenses are limited to a ten thousand dollar ($10,000) maximum per transplant.

(I) Lodging—maximum lodging expenses shall not exceed the per diem rates as established annually by U.S. General Services Administration (GSA) for a specific city or county. Go to www.gsa.gov for per diem rates.
(II) Travel—IRS standard medical mileage rates (same as flexible spending account (FSA) reimbursement).
(III) Meals—not covered.

B. Non-network. Charges above the maximum for services rendered at a non-network facility are the member’s responsibility and do not apply to the member’s deductible or out-of-pocket maximum. Travel, lodging, and meals are not covered;

46. Urgent care. Member encounter with a provider for urgent care is covered based on the service, procedure, or related treatment plan; and

47. Vision. One (1) routine exam and refraction is covered per calendar year.


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISMOUIR CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 3—Public Entity Membership

EMERGENCY AMENDMENT

22 CSR 10-3.058 PPO 750 Plan Benefit Provisions and Covered Charges. The Missouri Consolidated Health Care Plan is amending sections (5), (11), (12), and (14).

PURPOSE: This amendment adds one hundred percent (100%) coverage for sterilization procedure for men, revisions maximum plan payment, timely filing timeframe, and services performed in another country for the PPO 750 Plan.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (public entity employee members, retirees, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to public entity employee members, retirees, and their families as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(5) The following services are not subject to deductible, coinsurance, or copayment requirements and will be paid at one hundred percent (100%) when provided by a network provider:
(C) A newborn’s initial hospitalization until discharge or transfer to another facility if the mother is a Missouri Consolidated Health Care Plan (MCHCP) member at the time of birth;

(D) Four (4) Diabetes Self-Management Education visits\(\text{.}\) and

(E) Sterilization procedure for men.

(11) Maximum plan payment—non-network medical claims that are not otherwise subject to a contractual discount arrangement are processed at one hundred ten percent (110%) of Medicare reimbursement for non-network professional claims and following the claim administrator’s standard practice for non-network facility claims. Members may be held liable for the amount of the fee above the allowed amount.

(12) Any claim must be initially submitted within twelve (12) months following the date of service, unless other specified in the network provider contract. The plan reserves the right to deny claims not timely filed. A provider initiated correction to the originally filed claim must be submitted within the timeframe agreed in the provider contract, but not to exceed three hundred sixty-five (365) days from adjudication of the originally filed claim. Any claims reprocessed as primary based on action taken by Medicare or Medicaid must be initiated within three (3) years of the claim being incurred.

(14) Services performed in a country other than the United States may be covered if the service is included in 22 CSR 10-2055/3.057. Emergency and urgent care services are covered as a network benefit. All other non-emergency services are covered as a non-network benefit/determined by the claims administrator. If the service is provided by a non-network provider, the member may be required to provide payment to the provider and then file a claim for reimbursement subject to timely filing limits.

PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 3—Public Entity Membership

EMERGENCY AMENDMENT

22 CSR 10-3.059 PPO 1250 Plan Benefit Provisions and Covered Charges. The Missouri Consolidated Health Care Plan is amending sections (5), (11), (12), and (14).

PURPOSE: This amendment adds one hundred percent (100%) coverage for sterilization procedure for men, rewrites maximum plan payment, timely filing timeframe, and services performed in another country for the PPO 1250 Plan.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (public entity employee members, retirees, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to public entity employee members, retirees, and their families as one method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(5) The following services are not subject to deductible, coinsurance, or copayment requirements and will be paid at one hundred percent (100%) when provided by a network provider:

(A) Preventive care;

(B) Nutrition counseling;

(C) A newborn’s initial hospitalization until discharge or transfer to another facility if the mother is a Missouri Consolidated Health Care Plan (MCHCP) member at the time of birth;

(D) Four (4) Diabetes Self-Management Education visits\(\text{.}\)

(E) Sterilization procedure for men.

(11) Maximum plan payment—non-network medical claims that are not otherwise subject to a contractual discount arrangement are allowed at one hundred ten percent (110%) of Medicare reimbursement for non-network professional claims and following the claim administrator’s standard practice for non-network facility claims. Members may be held liable for the amount of the fee above the allowed amount.

(12) Any claim must be initially submitted within twelve (12) months following the date of service, unless other specified in the network provider contract. The plan reserves the right to deny claims not timely filed. A provider initiated correction to the originally filed claim must be submitted within the timeframe agreed in the provider contract, but not to exceed three hundred sixty-five (365) days from adjudication of the originally filed claim. Any claims reprocessed as primary based on action taken by Medicare or Medicaid must be initiated within three (3) years of the claim being incurred.

(14) Services performed in a country other than the United States may be covered if the service is included in 22 CSR 10-2055/3.057. Emergency and urgent care services are covered as a network benefit. All other non-emergency services are covered as a non-network benefit/determined by the claims administrator. If the service is provided by a non-network provider, the member may be required to provide payment to the provider and then file a claim for reimbursement subject to timely filing limits.

AUTHORITY: section 103.059, RSMo 2016. Emergency rule filed Oct. 31, 2018, effective Jan. 1, 2019, expired June 29, 2019. Original rule filed Oct. 30, 2018, effective May 30, 2019. Emergency amendment filed Oct. 30, 2019, effective Jan. 1, 2020, expires June 28, 2020. A proposed amendment covering this same material is published in this issue of the Missouri Register. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

AUTHORITY: section 103.059, RSMo 2016. Emergency rule filed Oct. 31, 2018, effective Jan. 1, 2019, expired June 29, 2019. Original rule filed Oct. 30, 2018, effective May 30, 2019. Emergency amendment filed Oct. 30, 2019, effective Jan. 1, 2020, expires June 28, 2020. A proposed amendment covering this same material is published in this issue of the Missouri Register. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.
PUBLIC COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Emergency Rules


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN

Division 10—Health Care Plan
Chapter 3—Public Entity Membership

EMERGENCY AMENDMENT

22 CSR 10-3.061 Plan Limitations. The Missouri Consolidated Health Care Plan is amending section (1).

PURPOSE: This rule establishes the policy of the board of trustees in regard to the PPO 750 Plan, PPO 1250 Plan, and Health Savings Account (HSA) Plan limitations of the Missouri Consolidated Health Care Plan.

PURPOSE: This amendment revises the following benefits not payable by the plan: alternative therapies, assistant surgeon services, birthing center, home births, nocturnal enuresis alarm, therapy, vaccinations requested by a third party, and numbers as necessary.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (public entity employee members, retirees, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to public entity employee members, retirees, and their families as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

1. Benefits shall not be payable for, or in connection with, any medical benefits, services, or supplies which do not come within the definition of covered charges. In addition, the items specified in this rule are not covered unless expressly stated otherwise and then only to the extent expressly provided herein or in 22 CSR 10-3.057 or 22 CSR 10-3.090.

2. Any state’s cash sickness or similar law, including any group insurance policy approved under such law.

3. (C) Alternative therapies—that are outside conventional medicine (including, but not limited to, acupuncture, acupressure, homeopathy, hypnosis, massage therapy, reflexology, and biofeedback) as determined by the claims administrator.

4. (E) Assistant surgeon services—unless determined to meet the clinical eligibility for coverage under the plan.

5. (F) (E) Athletic enhancement services and sports performance training.

6. (G) (F) Autopsy.

7. (H) Birthing center.

8. (I) (G) Blood donor expenses.

9. (J) (H) Blood pressure cuffs/monitors.

10. (K) (I) Care received without charge.

11. (L) (J) Charges exceeding the vendor contracted rate or benefit limit.

12. (M) (K) Charges resulting from the failure to appropriately cancel a scheduled appointment.

13. (N) (L) Childbirth classes.

14. (O) (M) Comfort and convenience items.

15. (P) (N) Cosmetic procedures.

16. (Q) (O) Custodial or domiciliary care—including services and supplies that assist members in the activities of daily living such as walking, getting in and out of bed, bathing, dressing, feeding, and using the toilet; preparation of special diets; supervision of medication that is usually self-administered; or other services that can be performed by persons who are not providers.

17. (R) (P) Dental care, including oral surgery.

18. (S) (Q) Devices or supplies bundled as part of a service are not separately covered.

19. (T) (R) Dialysis received through a non-network provider.

20. (U) (S) Educational or psychological testing unless part of a treatment program for covered services.

21. (V) (T) Examinations requested by a third party.

22. (W) (U) Exercise equipment.

23. (X) (V) Experimental/investigational/unproven services, procedures, supplies, or drugs as determined by the claims administrator.

24. (Y) (W) Eye services and associated expenses for orthoptics, eye exercises, radial keratotomy, LASIK, and other refractive eye surgery.

25. (Z) (X) Genetic testing based on family history alone, except for breast cancer susceptibility gene (BRCA) testing.

26. (AA) (Y) Health and athletic club membership—including costs of enrollment.

27. (BB) (Z) Hearing aid replacement batteries.

28. (CC) Home births.

29. (DD) (AA) Infertility treatment beyond the covered services to diagnose the condition.

30. (EE) (BB) Infusions received through a non-network provider.

31. (FF) (CC) Level of care, greater than is needed for the treatment of the illness or injury.

32. (GG) (DD) Long-term care.

33. (HH) (EE) Maxillofacial surgery.

34. (III) (FF) Medical care and supplies to the extent that they are payable under—

1. A plan or program operated by a national government or one (1) of its agencies; or

2. Any state’s cash sickness or similar law, including any group insurance policy approved under such law.

35. (JJJ) (GG) Medical service performed by a family member—including a person who ordinarily resides in the subscriber’s household or is related to the member, such as a spouse, parent, child, sibling, or brother/sister-in-law.

36. (KK) (HH) Military service-connected injury or illness—including expenses relating to Veterans Affairs or a military hospital.
Never events—never events on a list compiled by the National Quality Forum of inexcusable outcomes in a health care setting.

Nocturnal enuresis alarm.

Drugs that the pharmacy benefit manager (PBM) has excluded from the formulary and will not cover as a non-formulary drug unless it is approved in advance by the PBM.

Non-medically necessary services.

Non-provider allergy services or associated expenses relating to an allergic condition, including installation of air filters, air purifiers, or air ventilation system cleaning.

Non-reusable disposable supplies.

Online weight management programs.

Other charges as follows:

1. Charges that would not otherwise be incurred if the subscriber was not covered by the plan;
2. Charges for which the subscriber or his/her dependents are not legally obligated to pay including, but not limited to, any portion of any charges that are discounted;
3. Charges made in the subscriber’s name but which are actually due to the injury or illness of a different person not covered by the plan; and
4. No coverage for miscellaneous service charges including, but not limited to, charges for telephone consultations, administrative fees such as filling out paperwork or copy charges, or late payments.

Over-the-counter medications with or without a prescription including, but not limited to, analgesics, antipyretics, non-sedating antihistamines, unless otherwise covered as a preventive service.

Physical and recreational fitness.

Private-duty nursing.

Routine foot care without the presence of systemic disease that affects lower extremities.

Services obtained at a government facility if care is provided without charge.

Sex therapy.

Surrogacy—pregnancy coverage is limited to plan member.

Telehealth site origination fees or costs for the provision of telehealth services are not covered.

Physical, occupational, and speech therapy are not covered for the following:

1. Physical therapy—
   A. Treatment provided to prevent or slow deterioration in function or prevent reoccurrences;
   B. Treatment intended to improve or maintain general physical condition;
   C. Long-term rehabilitative services when significant therapeutic improvement is not expected;
   D. Physical therapy that duplicates services already being provided as part of an authorized therapy program through another therapy discipline (e.g., occupational therapy);
   E. Work hardening programs;
   F. Vocational rehabilitation programs and any programs with the primary goal of returning an individual to work;
   G. Group occupational therapy (because it is not one-on-one, individualized to the specific person’s needs); and
   H. Driving safety/driver training; and
2. Speech or voice therapy—
   A. Any computer-based learning program for speech or voice training purposes;
   B. School speech programs;
   C. Speech or voice therapy that duplicates services already being provided as part of an authorized therapy program through another therapy discipline (e.g., occupational therapy);
   D. Group speech or voice therapy (because it is not one-on-one, individualized to the specific person’s needs);
   E. Maintenance programs of routine, repetitive drills/exercises that do not require the skills of a speech-language therapist and that can be reinforced by the individual or caregiver;
   F. Vocational rehabilitation programs and any programs with the primary goal of returning an individual to work;
   G. Therapy or treatment provided to prevent or slow deterioration in function or prevent reoccurrences;
   H. Therapy or treatment provided to improve or enhance job, school, or recreational performance; and
   I. Long-term rehabilitative services when significant therapeutic improvement is not expected.

Travel expenses.

Vaccinations requested by third party.

Workers’ Compensation services or supplies for an illness or injury eligible for, or covered by, any federal, state, or local government Workers’ Compensation Act, occupational disease law, or other similar legislation.


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 3—Public Entity Membership

EMERGENCY AMENDMENT
22 CSR 10-3.070 Coordination of Benefits. The Missouri Consolidated Health Care Plan is amending section (3).

PURPOSE: This amendment revises the order of benefit determination rules and renumbers as necessary.
EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (public entity employees, retirees, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP), from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to public entity employee members, retirees, and their families as one (1) method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(3) Order of Benefit Determination Rules.

(B) Rules. MCHCP determines its order of benefits [using the first of the following rules which applies] as follows:

1. Active/inactive employee. The benefits of the plan which covers the person as an employee who is neither laid off nor retired (or as that employee’s dependent) are determined before those of the plan which covers that person as a laid off or retired employee (or as that employee’s dependent);

2. Nondependent/dependent. The benefits of the plan which covers the person as an employer or subscriber (that is, other than as a dependent) are determined before those of the plan which covers the person as a dependent;]

1. Non-Dependent/Dependent:

A. The plan which covers the member as an employee or subscriber is primary.

B. The plan which covers the member as dependent is secondary.

2. Active/layoff. The plan that covers the member or dependent through the member’s active employment is primary. A plan that covers the member or dependent through the member’s status as a laid off employee.

3. Retiree. The plan that covers the member or dependent through the member’s active employment is primary to a plan that covers the member or dependent through the member’s status as a retiree.

3.4. Medicare.

A. If a member is an active employee and has Medicare, MCHCP is the primary plan for the active employee and his/her dependents. Medicare is the secondary plan except for members with end stage renal disease (ESRD) as defined in subparagraph (3)(B)/3.4.C.

B. If a member is a retiree and has Medicare, Medicare is the primary plan for the retiree and his/her Medicare-eligible dependents. MCHCP is the secondary plan.

C. If a member or his/her dependents are eligible for Medicare solely because of ESRD, the member’s MCHCP plan is primary to Medicare during the first thirty (30) months of Medicare eligibility for home peritoneal dialysis or home hemodialysis and thirty-three (33) months for in-center dialysis. After the thirty (30) or thirty-three (33) months, Medicare becomes primary, and claims are submitted first to Medicare, then to MCHCP for secondary coverage. The member is responsible for notifying MCHCP of his/her Medicare status.

4.5. Dependent child/parents not separated or divorced.

When MCHCP and another plan cover the same child as a dependent of different [persons, called] parents—

A. The benefits of the plan of the parent whose birthday falls earlier in a year are determined before those of the plan of the parent whose birthday falls later in that year; but

B. If both parents have the same birthday, the benefits of the plan which covered one (1) parent longer are determined before those of the plans which covered the other parent for a shorter period of time;

5.6. Dependent child/separated or divorced, or never married.

If two (2) or more plans cover a person as a dependent child of divorced, separated, or never married parents, benefits for the child are determined in this order—

A. First, the plan of the parent with custody of the child;

B. Then, the plan of the spouse of the parent with the custody of the child;

C. Then, the plan of the parent not having custody of the child; and

D. Finally, the plan of the spouse of the parent not having custody of the child. However, if the specific terms of a court decree state that one (1) of the parents is responsible for the health care expense of the child and the entity obligated to pay or provide the benefits of the plan of that parent or spouse of the other parent has actual knowledge of those terms, the benefits of that plan are determined first. The plan of the other parent shall be the secondary plan. This paragraph does not apply with respect to any claim determination period or plan year during which any benefits are actually paid or provided before the entity has that actual knowledge;

6.7. Joint custody. If the specific terms of a court decree state that the parents shall share joint custody, without stating that one (1) of the parents is responsible for the health care expenses of the child, the plans covering the child shall follow the order of benefit determination rules outlined in paragraph (3)(B)/4.5.;

7.8. Dependent child/parents both parents covered by MCHCP. If both parents are covered by MCHCP and both parents cover the child as a dependent, MCHCP will not coordinate benefits with itself;

8.9. [The plan that covers the member as a spouse is primary over the plan that covers the member as a dependent child] When an adult dependent is covered by both spouse and parent, the benefits of the plan which covered a person longer are determined before those of the plan which covered that person for the shorter term; and

9.10. Longer/shorter length of coverage. If none of the previous rules determines the order of benefits, the benefits of the plan which covered a person longer are determined before those of the plan which covered that person for the shorter term.


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500)
PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.

Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 3—Public Entity Membership

EMERGENCY AMENDMENT

22 CSR 10-3.075 Review and Appeals Procedure. The Missouri Consolidated Health Care Plan is amending sections (1), (3), and (5).

PURPOSE: This amendment revises the claim submission and initial benefit determinations time frames, updates the name and appeal contact information for the third party administrator.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (public entity employee members, retirees, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to public entity employee members, retirees, and their families as one method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(1) Claims Submissions and Initial Benefit Determinations.

(A) Medical and pharmacy service claims are divided into three (3) types: pre-service, post-service, and concurrent claims.

1. Pre-service claims are requests for approval that the plan or vendor requires a member to obtain before getting medical care or filling a prescription, such as prior authorization or a decision whether a treatment, procedure, or medication is medically necessary.

A. Pre-service claims must be decided within a reasonable period of time appropriate to the medical circumstances, but no later than [fifteen (15) days] twenty (20) business days from the date the vendor receives the claim. The vendor may extend the time period up to an additional [fifteen (15)] twenty (20) days if, for reasons beyond the vendor’s control, the decision cannot be made within the first [fifteen (15)] twenty (20) days. The vendor must notify the member prior to the expiration of the first fifteen- (15-) day period, explain the reason for the delay, and request any additional information. If more information is requested, the member has at least forty-five (45) days to provide the information to the vendor. The vendor then must decide the claim no later than fifteen (15) days after the additional information is supplied or after the period of time allowed to supply it ends, whichever is first.

B. Urgent care claims are a special type of pre-service claim that require a quicker decision because waiting the standard time could seriously jeopardize the member’s life, health, or ability to regain maximum function. A request for an urgent care claim may be submitted verbally or in writing and will be decided within seventy-two (72) hours. Written confirmation of the decision will be sent by the vendor [as soon as possible thereafter] within three (3) business days.

2. Post-service claims must be decided within a reasonable period of time, not later than [thirty (30) days] twenty (20) business days after the vendor receives the claim. If, because of reasons beyond the vendor’s control, more time is needed to review the claim, the vendor may extend the time period up to an additional [fifteen (15)] thirty (30) days. The vendor must notify the member prior to the expiration of the first [fifteen- (15-) day] twenty (20-) day period, explain the reason for the delay, and request any additional information. If more information is requested, the member has at least forty-five (45) days to provide the information to the vendor. The vendor then must decide the claim no later than [fifteen (15) days] thirty (30) days after the additional information is supplied or after the period of time allowed to supply it ends, whichever is first.

3. Concurrent claims are claims related to an ongoing course of previously approved treatment. If the plan or vendor has approved an ongoing course of treatment to be provided over a period of time or number of treatments, any reduction or termination of the course of treatment will be treated as a benefit denial. The plan or vendor will notify a member in writing prior to reducing or ending a previously approved course of treatment in sufficient time to allow the member, or the member’s provider, to appeal and obtain a determination before the benefit is reduced or terminated.

B. Urgent care claims are a special type of pre-service claim that require a quicker decision because waiting the standard time could seriously jeopardize the member’s life, health, or ability to regain maximum function. A request for an urgent care claim may be submitted verbally or in writing and will be decided within seventy-two (72) hours. Written confirmation of the decision will be sent by the vendor [as soon as possible thereafter] within three (3) business days.

2. Post-service claims are all other claims for services including claims after medical or pharmacy services have been provided, such as requests for reimbursement or payment of the costs for the services provided.

A. Post-service claims must be decided within a reasonable period of time, not later than [thirty (30) days] twenty (20) business days after the vendor receives the claim. If, because of reasons beyond the vendor’s control, more time is needed to review the claim, the vendor may extend the time period up to an additional [fifteen (15)] thirty (30) days. The vendor must notify the member prior to the expiration of the first [fifteen- (15-) day] twenty (20-) day period, explain the reason for the delay, and request any additional information. If more information is requested, the member has at least forty-five (45) days to provide the information to the vendor. The vendor then must decide the claim no later than [fifteen (15) days] thirty (30) days after the additional information is supplied or after the period of time allowed to supply it ends, whichever is first.

3. Concurrent claims are claims related to an ongoing course of previously approved treatment. If the plan or vendor has approved an ongoing course of treatment to be provided over a period of time or number of treatments, any reduction or termination of the course of treatment will be treated as a benefit denial. The plan or vendor will notify a member in writing prior to reducing or ending a previously approved course of treatment in sufficient time to allow the member, or the member’s provider, to appeal and obtain a determination before the benefit is reduced or terminated.

(3) Appeal Process for Medical and Pharmacy Determinations.

(A) Definitions. Notwithstanding any other rule in this chapter to the contrary, for purposes of a member’s right to appeal any adverse benefit determination made by the plan, the plan administrator, a claims administrator, or a medical or pharmacy benefit vendor, relating to the provision of health care benefits, other than those provided in connection with the plan’s dental or vision benefit offering, the following definitions apply:

1. Adverse benefit determination. An adverse benefit determination means any of the following:

A. A denial, reduction, or termination of, or a failure to provide or make payment (in whole or in part) for a benefit, including any denial, reduction, termination, or failure to provide or make payment that is based on a determination of an individual’s eligibility to participate in the plan;

B. A denial, reduction, or termination of, or a failure to provide or make payment (in whole or in part) for a benefit resulting from the application of any utilization review, as well as a failure to cover an item or service for which benefits are otherwise provided because it is determined to be experimental or investigational or not medically necessary or appropriate; or

C. Any rescission of coverage after an individual has been covered under the plan;

2. Appeal (or internal appeal). An appeal or internal appeal means review by the plan, the plan administrator, a claims administrator, or a medical or pharmacy benefit vendor of an adverse benefit
determination;

3. Claimant. Claimant means an individual who makes a claim under this subsection. For purposes of this subsection, references to claimant include a claimant’s authorized representative.

4. External review. The United States Department of Health and Human Services (HHS) conducts external reviews for adverse benefit determinations regarding medical and pharmacy benefits administered by [UMR, Aetna, Anthem, and Express Scripts, Inc.] that involve medical judgment (including, but not limited to, those based on medical necessity, appropriateness, health care setting, level of care, or effectiveness of a covered benefit; or a determination that a treatment is experimental or investigational) and a rescission of coverage (regardless of whether or not the rescission has any effect on any particular benefit at that time);

5. Final internal adverse benefit determination. A final internal adverse benefit determination means an adverse benefit determination that has been upheld by the plan, the plan administrator, a claims administrator, or a medical or pharmacy benefit vendor at the completion of the internal appeals process under this subsection, or an adverse benefit determination with respect to which the internal appeals process has been deemed exhausted by application of applicable state or federal law;

6. Final external review decision. A final external review decision means a determination rendered under the external review process at the conclusion of an external review; and

7. Rescission. A rescission means a termination or discontinuance of medical or pharmacy coverage that has retroactive effect, except that a termination or discontinuance of coverage is not a rescission if–

(I) the termination or discontinuance of coverage has only a prospective effect; or

(A) Adverse benefit determinations denying or terminating an individual’s coverage under the plan based on a determination of the individual’s eligibility to participate in the plan or the failure to pay premiums, or any rescission of coverage based on fraud or intentional misrepresentation of a member or authorized representative of a member are appealable exclusively to the Missouri Consolidated Health Care Plan (MCHCP) Board of Trustees (board);

(B) If, because of reasons beyond the vendor’s control, more time is needed to review the appeal, the vendor may extend the time period up to an additional [fifteen (15)] thirty (30) days. The vendor must notify the member prior to the expiration of the first [fifteen- (15-) ] twenty- (20-) day period, explain the reason for the delay, and request any additional information. If more information is requested, the member has at least forty-five (45) days to provide the information to the vendor. The vendor then must decide the claim no later than [fifteen (15)] thirty (30) days after the additional information is supplied or after the period of time allowed to supply it ends, whichever is first. [Written confirmation of the decision will be sent by the vendor within fifteen (15) business days.]

(III) An expedited appeal of an adverse benefit determination may be requested when a decision is related to a pre-service claim for urgent care. Expedited appeals will be reviewed by the vendor by someone who was not involved in the original decision and will include consultation with a qualified medical professional if a medical judgment is involved. Expedited appeals will be responded to within seventy-two (72) hours after receiving a request for an expedited review with written confirmation of the decision to the member within thirty (30) working days of providing notification of the determination.

(IV) Second level appeals must be submitted in writing within sixty (60) days of the date of the first level appeal decision letter that upholds the original adverse benefit determination. Second level appeals should include any additional information or documentation to support the reason the member believes the first level appeal decision should be overturned. Second level appeals will be reviewed by the vendor by someone who was not involved in the original decision or first level appeal and will include consultation with a qualified medical professional if a medical judgment is involved. Second level medical appeals shall be responded to in writing to the member within thirty (30) days for post-service claims and within fifteen (15) days for pre-service claims from the date the vendor received the second level appeal request.

(A) if, because of reasons beyond the vendor’s control, more time is needed to review the appeal, the vendor may extend the time period up to an additional fifteen (15) days. The vendor must notify the member prior to the expiration of the first fifteen- (15-) day period, explain the reason for the delay, and request any additional information. If more information is requested, the member has at least forty-five (45) days to provide the information to the vendor. The vendor then must decide the claim no later than fifteen (15) days after the additional information is supplied or after the period of time allowed to supply it ends, whichever is first.

(V) For members with medical coverage through UMR—

(a) First and second level pre-service, first and second level post-service, and concurrent claim appeals must be submitted in writing to—

Anthem Blue Cross and Blue Shield
Attn: Grievance Department
PO Box 105568
Atlanta, Georgia 30348-5568
or by fax to (800) 859-3046
(b) First and second level post-service appeals must be sent in writing to—
UMR Claims Appeal Unit
PO Box 30546
Salt Lake City, UT 84130-0546
or by fax to (877) 291-3248.

(c)(b) Expedited pre-service appeals must may be communicated submitted by calling (800) 808-4424, ext. 15227/ (877) 333-7488 or by submitting a written fax to (877) 615-6584, Attention: Appeals Unit (800) 368-3238.

C. The internal review process for adverse benefit determinations relating to pharmacy and the Pharmacy Lock-In Program consists of one (1) level of internal review provided by the pharmacy vendor.

(I) Pharmacy appeals. Pharmacy appeals and Pharmacy Lock-In Program appeals must identify the matter being appealed and should include the member’s (and dependent’s, if applicable) name, the date the member attempted to fill the prescription, the prescribing physician’s name, the drug name and quantity, the cost of the prescription, if applicable, and any applicable reason(s) relevant to the appeal including: the reason(s) the member believes the claim should be paid, the reason(s) the member believes the claim should not be included in the Pharmacy Lock-In Program, and any other written documentation to support the member’s belief that the original decision should be overturned.

(II) All pharmacy appeals must be submitted in writing to—
Express Scripts
Attn: Clinical Appeals Department
PO Box 66588
St. Louis, MO 63116-6588
or by fax to (877) 852-4070

(III) All Pharmacy Lock-In Program appeals must be submitted in writing to—
Express Scripts
Drug Utilization Review Program
Mail Stop HQ3W03
One Express Way
St. Louis, MO 63121

(iv) Pharmacy appeals will be reviewed by someone who was not involved in the original decision and the reviewer will consult with a qualified medical professional if a medical judgment is involved. Pharmacy appeals will be responded to in writing to the member within sixty (60) days for post-service claims and thirty (30) days for pre-service claims from the date the vendor received the appeal request.

(V) The Pharmacy Benefit Manager will respond to Pharmacy Lock-In Program appeals in writing to the member within thirty (30) days from the date the Pharmacy Benefit Manager received the appeal request.

D. Members may seek external review only after they have exhausted all applicable levels of internal review or received a final internal adverse benefit determination.

(I) A claimant or authorized representative may file a written request for an external review within four (4) months after the date of receipt of a final internal adverse benefit determination.

(II) The claimant can submit an external review request in writing to—
HHS Federal Request
MAXIMUS Federal Services

3750 Monroe Ave., Suite 705
Pittsford, NY 14534
or by fax to (888) 866-6190
or to request a review online at http://www.externalappeal.com/

(III) The claimant may call the toll-free number (888) 866-6205 with any questions or concerns during the external review process and can submit additional written comments to the external reviewer at the mailing address above.

(IV) The external review decision will be made as expeditiously as possible and within forty-five (45) days after receipt of the request for the external review.

(V) A claimant may make a written or oral request for an expedited external review if the adverse benefit determination involves a medical condition of the claimant for which the timeframe for completion of a standard external review would seriously jeopardize the life or health of the claimant; or would jeopardize the claimant’s ability to regain maximum function; or if the final internal adverse benefit determination involves an admission, availability of care, continued stay, or health care item or service for which the claimant received services, but has not been discharged from a facility.

3. For all internal appeals of adverse benefit determinations, the plan or the vendor reviewing the appeal will provide the member, free of charge, with any new or additional evidence or rationale considered, relied upon, or generated by the plan or the vendor in connection with reviewing the claim or the appeal and will give the member an opportunity to respond to such new evidence or rationale before issuing a final internal adverse determination.

5. In reviewing appeals, notwithstanding any other rule, the board and/or staff may grant any appeals when there is credible evidence to support approval under the following guidelines:

(I) MCHCP may approve an appeal regarding plan changes retrospectively for subscribers who are new employees within thirty (30) days of election of coverage if no claims have been filed with the previous carrier. If a subscriber has his/her premium collected pre-tax by qualified payroll deduction through a cafeteria plan, changes may be approved if the reason given is allowed by the cafeteria plan; and

(J) Once per lifetime of the account, MCHCP may approve an appeal where a subscriber missed a deadline. MCHCP may only approve an appeal under this guideline if the appeal is received within sixty (60) days of the missed deadline. This guideline may not be used to approve an appeal of a voluntary cancellation or an appeal of a deadline that is statutorily mandated; and,

[(K) MCHCP may approve an appeal to change a subscriber’s medical plan vendor prospectively, once per lifetime of the account. This appeal guideline may not be used for a subscriber to change the type of medical plan design elected during open enrollment.]


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.
Title 22—MISSOURI CONSOLIDATED HEALTH CARE PLAN
Division 10—Health Care Plan
Chapter 3—Public Entity Membership

EMERGENCY AMENDMENT

22 CSR 10-3.090 Pharmacy Benefit Summary. The Missouri Consolidated Health Care Plan is amending section (1).

PURPOSE: This amendment adds coinsurance dollar limits for the Health Savings Account (HSA) Plan, revises services covered at one hundred percent (100%), and renumbers as necessary.

EMERGENCY STATEMENT: This emergency amendment must be in place by January 1, 2020, in accordance with the new plan year. Therefore, this emergency amendment is necessary to serve a compelling governmental interest of protecting members (public entity employee members, retirees, and their families) enrolled in the Missouri Consolidated Health Care Plan (MCHCP) from the unintended consequences of confusion regarding eligibility or availability of benefits and will allow members to take advantage of opportunities for reduced premiums for more affordable options without which they may forgo coverage. Further, it clarifies member eligibility and responsibility for various types of eligible charges, beginning with the first day of coverage for the new plan year. It may also help ensure that inappropriate claims are not made against the state and help protect the MCHCP and its members from being subjected to unexpected and significant financial liability and/or litigation. It is imperative that this amendment be filed as an emergency amendment to maintain the integrity of the current health care plan. This emergency amendment fulfills the compelling governmental interest of offering access to more convenient and affordable medical services to public entity employee members, retirees, and their families as one method of protecting the MCHCP trust fund from more costly expenses. This emergency amendment reflects changes made to the plan by the Missouri Consolidated Health Care Plan Board of Trustees. A proposed amendment, which covers the same material, is published in this issue of the Missouri Register. This emergency amendment complies with the protections extended by the Missouri and United States Constitutions and limits its scope to the circumstances creating the emergency. The MCHCP follows procedures best calculated to assure fairness to all interested persons and parties under the circumstances. This emergency amendment was filed October 30, 2019, becomes effective January 1, 2020, and expires June 28, 2020.

(1) The pharmacy benefit provides coverage for prescription drugs. Vitamin and nutrient coverage is limited to prenatal agents, therapeutic agents for specific deficiencies and conditions, and hematopoietic agents as prescribed by a provider.

(A) PPO 750 Plan and PPO 1250 Plan Prescription Drug Coverage

1. Network

A. Preferred formulary generic drug copayment: Ten Dollars ($10) for up to a thirty-one- (31-) day supply; twenty dollars ($20) for up to a sixty- (60-) day supply; and thirty dollars ($30) for up to a ninety- (90-) day supply for a generic drug on the formulary; formulary generic birth control and tobacco cessation prescriptions covered at one hundred percent (100%).

B. Preferred formulary brand drug copayment: Forty dollars ($40) for up to a thirty-one- (31-) day supply; eighty dollars ($80) for up to a sixty- (60-) day supply; and one hundred twenty dollars ($120) for up to a ninety- (90-) day supply for a brand drug on the formulary; formulary brand birth control and tobacco cessation prescriptions covered at one hundred percent (100%).

C. Non-preferred formulary drug and approved excluded drug copayment: One hundred dollars ($100) for up to a thirty-one- (31-) day supply; two hundred dollars ($200) for up to a sixty- (60-) day supply; and three hundred dollars ($300) for up to a ninety- (90-) day supply for a drug not on the formulary.

D. Specialty drug (as designated as such by the PBM) copayment: Seventy-five dollars ($75) for up to a thirty-one- (31-) day supply for a specialty drug on the formulary.

E. Diabetic drug (as designated as such by the PBM) copayment: Fifty percent (50%) of the applicable network copayment.

F. Home delivery programs.

(I) Maintenance prescriptions may be filled through the pharmacy benefit manager’s (PBM’s) home delivery program. A member must choose how maintenance prescription(s) will be filled by notifying the PBM of his/her decision to fill a maintenance prescription through home delivery or retail pharmacy.

(a) If the member chooses to fill his/her maintenance prescription at a retail pharmacy and the member does not notify the PBM of his/her decision, the first two (2) maintenance prescription orders may be filled at the retail pharmacy. After the first two (2) orders are filled at the retail pharmacy, the member must notify the PBM of his/her decision to continue to fill the maintenance prescription at the retail pharmacy. If a member does not make a decision after the first two (2) orders are filled at the retail pharmacy, s/he will be charged the full discounted cost of the drug until the PBM has been notified of the decision and the amount charged will not apply to the out-of-pocket maximum.

(b) Once a member makes his/her delivery decision, the member can modify the decision by contacting the PBM.

(II) Specialty drugs are covered only through the specialty home delivery network for up to a thirty-one- (31-) day supply unless the PBM has determined that the specialty drug is eligible for up to a ninety- (90-) day supply. All specialty prescriptions must be filled through the PBM’s specialty pharmacy, unless the prescription is identified by the PBM as emergent. The first fill of a specialty prescription may be filled through a retail pharmacy.

(a) Specialty split-fill program—The specialty split-fill program applies to select specialty drugs as determined by the PBM. For the first three (3) months, members will be shipped a fifteen-(15-) day supply with a prorated copayment. If the member is able to continue with the medication, the remaining supply will be shipped with the remaining portion of the copayment. Starting with the fourth month, an up to thirty-one- (31-) day supply will be shipped if the member continues on treatment.

(III) Prescriptions filled through home delivery programs have the following copayments:

(a) Preferred formulary generic drug copayments: Ten dollars ($10) for up to a thirty-one- (31-) day supply; twenty dollars ($20) for up to a sixty- (60-) day supply; and twenty-five dollars ($25) for up to a ninety- (90-) day supply for a generic drug on the formulary.

(b) Preferred formulary brand drug copayments: Forty dollars ($40) for up to a thirty-one- (31-) day supply; eighty dollars ($80) for up to a sixty- (60-) day supply; and one hundred dollars ($100) for up to a ninety- (90-) day supply for a brand drug on the formulary.

(c) Non-preferred formulary drug and approved excluded drug copayments: One hundred dollars ($100) for up to a thirty-one- (31-) day supply; two hundred dollars ($200) for up to a sixty- (60-) day supply; and two hundred fifty dollars ($250) for up to a ninety- (90-) day supply for a drug not on the formulary.

(d) Specialty drug (as designated as such by the PBM) copayment: Seventy-five dollars ($75) for up to a thirty-one- (31-) day supply for a specialty drug on the formulary.

G. Diabetic drug (as designated as such by the PBM) copayment: Fifty percent (50%) of the applicable network copayment.

H. Only one (1) copayment is charged if a combination of different manufactured dosage amounts must be dispensed in order to fill a prescribed single dosage amount.
I. The copayment for a compound drug is based on the primary drug in the compound. The primary drug in a compound is the most expensive prescription drug in the mix. If any ingredient in the compound is excluded by the plan, the compound will be denied.

J. If the copayment amount is more than the cost of the drug, the member is only responsible for the cost of the drug.

K. If the physician allows for generic substitution and the member chooses a brand-name drug, the member is responsible for the generic copayment and the cost difference between the brand-name and generic drug which shall not apply to the out-of-pocket maximum.

L. Preferred select brand drugs, as determined by the PBM:

Ten dollars ($10) for up to a thirty-one- (31-) day supply; twenty dollars ($20) for up to a sixty- (60-) day supply; and twenty-five dollars ($25) for up to a ninety- (90-) day supply.

M. Prescription drugs and prescribed over-the-counter drugs as recommended by the U.S. Preventive Services Task Force (categories A and B) and, for women, by the Health Resources and Services Administration are covered at one hundred percent (100%) when filled at a network pharmacy. The following are also covered at one hundred percent (100%) when filled at a network pharmacy:

(I) Vaccine recommended by the Advisory Committee on Immunization Practices of the Centers for Disease Control and Prevention;

II. (III) Generic Tamoxifen, generic Raloxifene, and brand Soltamox for prevention of breast cancer;

II. (II) Prescribed preferred diabetic test strips and lancets; and

II. (III) One (1) preferred glucometer.

2. Non-network: If a member chooses to use a non-network pharmacy for non-specialty prescriptions, s/he will be required to pay the full cost of the prescription and then file a claim with the PBM. The PBM will reimburse the cost of the drug based on the network discounted amount as determined by the PBM, less the applicable network copayment.


A. Network and non-network out-of-pocket maximums are separate.

B. The family out-of-pocket maximum is an aggregate of applicable charges received by all covered family members of the plan. Any combination of covered family member applicable charges may be used to meet the family out-of-pocket maximum. Applicable charges received by one (1) family member may only meet the individual out-of-pocket maximum amount.

C. Network individual—four thousand one hundred fifty dollars ($4,150).

D. Network family—eight thousand three hundred dollars ($8,300).

E. Non-network—no maximum.

(B) Health Savings Account (HSA) Plan Prescription Drug Coverage. Medical and pharmacy expenses are combined to apply toward the appropriate network or non-network deductible and out-of-pocket maximum specified in 22 CSR 10-3.055.

1. Network.

A. Preferred formulary generic drug: Ten percent (10%) coinsurance up to fifty dollars ($50) per thirty-one- (31-) day supply after deductible has been met for a generic drug on the formulary.

B. Preferred formulary brand drug: Twenty percent (20%) coinsurance up to one hundred dollars ($100) per thirty-one- (31-) day supply after deductible has been met for a brand drug on the formulary.

C. Non-preferred formulary drug and approved excluded drug: Forty percent (40%) coinsurance after deductible has been met.

D. Diabetic drug (as designated as such by the PBM) coinsurance: fifty percent (50%) of the applicable network coinsurance after deductible has been met, not to exceed:

(I) Twenty-five dollars ($25) per thirty-one- (31-) day supply for generic drugs;

(II) Fifty dollars ($50) per thirty-one- (31-) day supply for preferred formulary brand drug; and

(III) One hundred dollars ($100) per thirty-one- (31-) day supply for non-preferred formulary drug.

E. Home delivery program.

(I) Maintenance prescriptions may be filled through the PBM’s home delivery program. A member must choose how maintenance prescriptions will be filled by notifying the PBM of his/her decision to fill a maintenance prescription through home delivery or retail pharmacy.

(a) If the member chooses to fill his/her maintenance prescription at a retail pharmacy and the member does not notify the PBM of his/her decision, the first two (2) maintenance prescription orders may be filled by the retail pharmacy. After the first two (2) orders are filled at the retail pharmacy, the member must notify the PBM of his/her decision to continue to fill the maintenance prescription at the retail pharmacy. If a member does not make a decision after the first two (2) orders are filled at the retail pharmacy, s/he will be charged the full discounted cost of the drug until the PBM has been notified of the decision.

(b) Once a member makes his/her delivery decision, the member can modify the decision by contacting the PBM.

(II) Specialty drugs are covered only through network home delivery for up to a thirty-one- (31-) day supply unless the PBM has determined that the specialty drug is eligible for up to a ninety- (90-) day supply. All specialty prescriptions must be filled through the PBM’s specialty pharmacy, unless the prescription is identified by the PBM as emergent. The first fill of a specialty prescription identified to be emergent, may be filled through a retail pharmacy.

(a) Specialty split-fill program—The specialty split-fill program applies to select specialty drugs as determined by the PBM. For the first three (3) months, members will be shipped a fifteen- (15-) day supply. If the member is able to continue with the medication, the remaining supply will be shipped. Starting with the fourth month, an up to thirty-one- (31-) day supply will be shipped if the member continues on treatment.

F. Prescription drugs and prescribed over-the-counter drugs as recommended by the U.S. Preventive Services Task Force (categories A and B) and, for women, by the Health Resources and Services Administration are covered at one hundred percent (100%) when filled at a network pharmacy. The following are also covered at one hundred percent (100%) when filled at a network pharmacy:

III. (G.) Vaccines and administration as recommended by the Advisory Committee on Immunization Practices of the Centers for Disease Control and Prevention; and

III. (H.) One (1) preferred glucometer.

III. (I) Maintenance prescriptions may be filled through the PBM’s home delivery program. A member must choose how maintenance prescriptions will be filled by notifying the PBM of his/her decision to fill a maintenance prescription through home delivery or retail pharmacy.

(a) If the member chooses to fill his/her maintenance prescription at a retail pharmacy and the member does not notify the PBM of his/her decision, the first two (2) maintenance prescription orders may be filled by the retail pharmacy. After the first two (2) orders are filled at the retail pharmacy, the member must notify the PBM of his/her decision to continue to fill the maintenance prescription at the retail pharmacy. If a member does not make a decision after the first two (2) orders are filled at the retail pharmacy, s/he will be charged the full discounted cost of the drug until the PBM has been notified of the decision.

(b) Once a member makes his/her delivery decision, the member can modify the decision by contacting the PBM.

(II) Specialty drugs are covered only through network home delivery for up to a thirty-one- (31-) day supply unless the PBM has determined that the specialty drug is eligible for up to a ninety- (90-) day supply. All specialty prescriptions must be filled through the PBM’s specialty pharmacy, unless the prescription is identified by the PBM as emergent. The first fill of a specialty prescription identified to be emergent, may be filled through a retail pharmacy.

(a) Specialty split-fill program—The specialty split-fill program applies to select specialty drugs as determined by the PBM. For the first three (3) months, members will be shipped a fifteen- (15-) day supply. If the member is able to continue with the medication, the remaining supply will be shipped. Starting with the fourth month, an up to thirty-one- (31-) day supply will be shipped if the member continues on treatment.
B. Preferred formulary brand drug: Forty percent (40%) coinsurance after deductible has been met for up to a thirty-one-(31-) day supply for a brand drug on the formulary.

C. Non-preferred formulary drug and approved excluded drug: Fifty percent (50%) coinsurance after deductible has been met for up to a thirty-one-(31-) day supply for a drug not on the formulary.

D. Diabetic drug (as designated by the PBM) coinsurance: Fifty percent (50%) of the applicable non-network coinsurance after deductible has been met.


PUBLIC COST: This emergency amendment will not cost state agencies or political subdivisions more than five hundred dollars ($500) in the aggregate.

PRIVATE COST: This emergency amendment will not cost private entities more than five hundred dollars ($500) in the aggregate.