
13. QUALITY MANAGEMENT

A. Quality Studies Medical Records Access

APPLIES TO:

A. This policy applies to all IEHP DualChoice Providers, Delegates and Hospitals.

POLICY:

A. For the purposes of medical data processing, quality of care assessment and other research, all Providers, Delegates and Hospitals must provide IEHP access to Members' medical records, at no cost to the Health Plan.^{1,2}

DEFINITION:

A. Delegate – For the purpose of this policy, this is defined as a medical group, Health Plans, IPA, or any contracted organization delegated to maintain and/or provide Member medical record access for use in quality studies.

PROCEDURES:

A. Quality Studies

1. IEHP performs quality studies to meet requirements of the California Department of Health Care Services (DHCS), Centers for Medicare and Medicaid Services (CMS) and the National Committee for Quality Assurance (NCQA). These studies cross over total IEHP Membership.
2. IEHP utilizes NCQA's Healthcare Effectiveness Data Information Set (HEDIS®) methodology for all applicable quality studies. For studies not addressed by HEDIS®, IEHP utilizes a format approved by the agency requesting the study.
3. To complete these studies according to required methodologies, IEHP must gather information both from administrative data (i.e., encounter data) and Members' medical records.

B. Delegate Pre-notification

1. IEHP notifies Delegates at least five (5) business days before Providers are contacted for medical record information.
2. Notification includes a description of the study purpose and requirements.

C. Provider and Hospital Notification

1. IEHP notifies Providers and Hospitals if any of their Members have been selected for inclusion in a quality study.
2. Notification includes a description of the study purpose and requirements as well as a list

¹ California Civil Code (Civ. Code) § 56.10

² Title 22, California Code of Regulations (CCR) § 51009

13. QUALITY MANAGEMENT

A. Quality Studies Medical Records Access

of the Members whose records are needed and the method of data collection.

3. IEHP collects medical record data in one of the following ways, depending on the nature of the study and the location of the Provider's office or Hospital:
 - a. IEHP staff may make appointments with the Provider's office or Hospital to visit the site for the purpose of medical record review and/or data collection. Data collection includes making photocopies and/or scanning hard copy medical records or downloading selected electronic medical records for study purposes.
 - b. IEHP may request that the Provider's office or Hospital retrieve the requested records and mail, fax or email records to IEHP.

D. Confidentiality

1. IEHP maintains compliance with the Health Information Portability and Accountability Act (HIPAA) requirements with all Member medical record information, including information used for the purpose of a quality study.
2. IEHP maintains strict confidentiality when using Member records for quality studies.
3. Members' identities are not disclosed in quality study results.
4. Abstracted data is archived and saved for a period of time determined by the study on an IEHP secure server.

INLAND EMPIRE HEALTH PLAN		
Chief Approval: <i>Signature on file</i>	Original Effective Date:	January 1, 2007
Chief Title: Chief Medical Officer	Revision Date:	January 1, 2023

13. QUALITY MANAGEMENT

B. Quality Management Program Overview for Members and Providers

APPLIES TO:

- A. This policy applies to all IEHP DualChoice Members and Providers.

POLICY:

- A. IEHP makes information about the Quality Management (QM) Program, including information on achieving established quality goals, available to all Members and Providers to keep them informed of quality improvement and quality management activities and outcomes.

PROCEDURES:

- A. IEHP provides the following information regarding the QM Program to Members and Providers via the IEHP website at www.iehp.org. QM Program information includes:
1. The “IEHP Annual Evaluation of Quality Management Executive Summary” addresses progress in achieving quality goals and contains yearly Healthcare Effectiveness Data and Information Set (HEDIS[®]) and Consumer Assessment of Healthcare Providers and Systems (CAHPS[®]) results.
 2. The “Quality Management Program Description” provides information on goals and objectives, QM and Quality Improvement (QI) activities addressing access to care, experience surveys, clinical practice guidelines and IEHP monitoring activities.
- B. Members or Providers who are not able to access the website or prefer a printed version of the QM Program information can request it through the following:
1. Calling the IEHP Member Services Department at (800) 440-IEHP (4347), or (800) 718-4347 for TTY users.
 2. Submitting a written request to IEHP at:

Inland Empire Health Plan
P.O. Box 1800
Rancho Cucamonga, CA 91729-1800
Attention: Quality Management Department

Upon receipt of a written request for information letter, the QM Department staff mails a packet to the requesting party consisting of the QM Program information.

- C. At least annually, IEHP reviews the results of the Quality Withhold measures in the IEHP Quality Improvement Subcommittee (QISC) that has representation from various departments across the organization. A brief summary of the results is included in the QM Program information.

13. QUALITY MANAGEMENT

B. Quality Management Program Overview for Members and Providers

- D. Members and Providers are advised to contact IEHP in writing if they have suggestions or would like further information on the QM/QI Program and activities.

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Chief Approval: <i>Signature on file</i>	Original Effective Date:	July 1, 2012
Chief Title: Chief Medical Officer	Revision Date:	January 1, 2023

13. QUALITY MANAGEMENT

C. Chaperone Guidance

APPLIES TO:

- A. This policy applies to all IEHP DualChoice Members and Providers.

POLICY:

- A. IEHP and its IPAs ensure Providers adopt a policy that Members are free to request a chaperone and ensure that the policy is communicated to Members.¹
- B. The presence of a chaperone during a clinical examination and treatment must be the clearly expressed choice of a Member; however, the default position should be that all intimate examinations are chaperoned.

PURPOSE:

- A. To ensure respect for the Member's dignity by providing a comfortable and considerate atmosphere for both the Member and the Provider.²

DEFINITION:

- A. Chaperone – A member of the Provider's medical staff whose job is to enhance the patient's and Provider's comfort, safety, privacy, security and dignity during sensitive exams or procedures.

PROCEDURES:

Provider's Responsibilities

- A. Providers should always honor a Member's request to have a chaperone.³
- B. Providers should in general, use a chaperone even when a patient's trusted companion is present.⁴
- C. Providers should ensure that all Members are offered a chaperone during any consultation, or intimate examination or procedure.
- D. The Member has the right to decline any chaperone offered. It is important to record in the Member's medical record that the offer was made, and that the Member declined
- E. Providers should provide the opportunity for private conversations with the Member without the chaperone present. Providers should minimize inquiries or history taking of a sensitive nature during a chaperoned examination.⁵

¹ American Medical Association (AMA), Code of Medical Ethics Opinion 1.2.4, Use of Chaperones, 04/26/2021

² Ibid.

³ Ibid.

⁴ Ibid.

⁵ Ibid.

13. QUALITY MANAGEMENT

C. Chaperone Guidance

- F. In instances, where the Member requests or accepts the offer of a chaperoned visit and one is not available at that time:
1. The Member must be given the opportunity to reschedule their appointment within a reasonable timeframe.
 2. Providers should contact the Member upon notice of unavailability of a chaperone, when necessary.
 3. If the seriousness of the condition would dictate that a delay is inappropriate, then this should be explained to the Member and recorded in their medical record.
 4. It is acceptable for the Provider (or other appropriate member of the clinical team) to perform an intimate examination without a chaperone if the situation is life-threatening or speed is essential in the care or treatment of the Member. This should be recorded in the Member's medical record.

Chaperone's Responsibilities

- A. Chaperoning should not be undertaken by anyone other than medical staff. This applies to all healthcare professionals working within a clinical or medical office setting.
- B. No family member or friend of a Member may be routinely expected to undertake any formal chaperoning role in normal circumstances.
- C. The chaperone must sign the Member's medical record indicating their presence during the visit.
- D. A chaperone who identifies unusual or unacceptable behavior by the Provider is expected to report to the: IEHP Compliance Hotline (866) 355-9038 or the California Medical Board at (800) 633-2322.

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Chief Approval: <i>Signature on file</i>	Original Effective date:	July 1, 2014
Chief Title: Chief Medical Officer	Revised Date:	January 1, 2023

13. QUALITY MANAGEMENT

D. Reporting Requirements Related to Provider Preventable Conditions

APPLIES TO:

- A. This policy applies to all IEHP network Providers who treat IEHP DualChoice Members.

POLICY:

- A. IEHP and its IPAs ensure Provider-Preventable Conditions (PPC) are reported to the California Department of Health Care Services (DHCS).^{1,2}

DEFINITIONS:

- A. Provider-Preventable Conditions (PPC) - These include both Category One (1) – Health Care-Acquired Conditions (HCACs) for any inpatient hospital setting and Category Two (2) – Other Provider Preventable Conditions (OPPC) for any health care setting (See Attachment, “Provider Preventable Conditions” in Section 13).³

PROCEDURES:

- A. All Providers must report PPCs through DHCS’ secure online reporting portal, which is found at <http://apps.dhcs.ca.gov/PPC/SecurityCode.aspx>.^{4,5}
- B. On a monthly basis, IEHP reviews encounter data submitted by network Providers for evidence of PPCs. IEHP’s HealthCare Informatics (HCI) team systematically screens and identifies potential PPCs among encounter data and ensure that confirmed PPCs are reported to DHCS Audits & Investigation (A&I) Division through its secure online reporting portal.⁶ Potential PPC identified through the monthly encounter data mining are processed as a Potential Quality Incident (PQI) case.
- C. Annually, IEHP reminds all network Providers of reporting requirements related to PPCs.⁷
- D. All network Providers are responsible for providing IEHP with a copy of all PPCs submitted to DHCS. Copies must be sent by fax at (909) 890-5545 within five (5) business days of reporting to DHCS.
- E. All completed and submitted PPC submissions are retained by IEHP.⁸

¹ Title 42, Code of Federal Regulations (CFR) § 438.3

² Department of Health Care Services (DHCS) Duals Plan Letter (DPL) 17-002 Supersedes DPL 15-002, “Reporting Requirements Related to Provider Preventable Conditions”

³ DHCS DPL 17-002

⁴ Ibid.

⁵ 42 CFR § 438.3

⁶ DHCS DPL 17-002

⁷ Ibid.

⁸ Ibid.

13. QUALITY MANAGEMENT

D. Reporting Requirements Related to Provider Preventable Conditions

F. IEHP reserves the right to recover or recoup any claim related to a PPC.⁹

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Chief Approval: <i>Signature on File</i>	Original Effective Date:	January 1, 2017
Chief Title: Chief Medical Officer	Revision Date:	January 1, 2023

⁹ 42 CFR § 438.3

13. QUALITY MANAGEMENT

E. Chronic Care Improvement Program (CCIP)

APPLIES TO:

- A. This policy applies to all IEHP DualChoice (HMO D-SNP) delegated IPAs and their assigned Members.

POLICY:

- A. IEHP and its IPAs are required to have an ongoing Chronic Care Improvement Program (CCIP) that serves the most vulnerable population identified by IEHP.¹

DEFINITIONS:

- A. Most Vulnerable population – IEHP has mechanisms in place to monitor and stratify its current dual-eligible population into risk categories that allow identification of the most vulnerable Members. IEHP provides details on their most vulnerable population and D-SNP Model of Care (MOC) via the IEHP website at www.iehp.org.

PURPOSE:

- A. The CCIP is intended to achieve the following objectives:²
1. To promote effective chronic disease management and the improvement of care and health outcomes for Members with chronic conditions.
 2. Support the IEHP focused population outlined in the Medicare D-SNP MOC;
 3. Include interventions that are above and beyond the Delegate’s inherent care coordination role and overall management of enrollees;
 4. Engage enrollees as partners in their care;
 5. Increase disease management and preventive services utilization;
 6. Improve health outcomes;
 7. Facilitate the development of targeted goals and specific interventions, aimed to improve one of the identified targeted conditions;
 8. Guard against potential health disparities; and
 9. Produce best practices.

PROCEDURES:

- A. IEHP and its IPAs must develop a methodology to identify Members with multiple or severe

¹ Title 42 Code of Federal Regulations (CFR) § 422.152

² Centers for Medicare and Medicaid Services (CMS) – Medicare Advantage (MA) – Chronic Care Improvement Program (CCIP) Resource Document (Updated 2020)

13. QUALITY MANAGEMENT

E. Chronic Care Improvement Program (CCIP)

chronic conditions who would benefit from participating in the CCIP (or Providers of such Members) as the target population.³

1. The conditions in the CCIP must be appropriate to meet the needs of its Membership and be tied to its most vulnerable population.
 2. IEHP and its IPAs are required to conduct the CCIP over a three-year (3) period.
- B. IEHP and its IPAs must conduct a comprehensive analysis of their target population to develop meaningful CCIP interventions. The non-exhaustive list below includes examples of intervention types that should be included in CCIPs:⁴
1. Care coordination to ensure Members receive care according to accepted standards of practice (i.e., clinical guidelines);
 2. Promotion of lifestyle changes and use of preventive services to slow the progression of the disease and/or prevent the development of complications and comorbidities;
 3. Effective disease management programs;
 4. Outreach to establish partnerships/collaboration with Providers, community groups, and stakeholders to leverage resources;
 5. Effective communication across the care continuum; and
 6. Education and outreach interventions to engage Members and caregivers as partners in care.
- C. IEHP and its IPAs must assess and internally document activities related to these quality initiatives on an ongoing basis, as well as modify interventions and/or processes as necessary.⁵
- D. IEHP and its IPAs must follow the Plan, Do, Study, Act (PDSA) quality improvement model as the overall structure for implementation and monitoring of the CCIP (see Attachment, “Chronic Care Improvement Program (CCIP) Planning & Reporting Document” in Section 13).⁶
- E. IEHP and its IPAs must comply with the CCIPs requirement of two (2) submissions per annual cycle (see Attachment, “Medicare Provider Reporting Requirements Schedule” in Section 25).⁷
1. CCIP Program Launch Submission – Due as the first (1st) semi-annual submission.
 - a. The CCIP program launch submission includes two (2) components:
 - 1) The CCIP Overview

³ 42 CFR § 422.152(c)(1)(i)

⁴ CMS MA CCIP Resource Document (Updated 2020)

⁵ Ibid.

⁶ Ibid.

⁷ 42 CFR § 422.152(c)

13. QUALITY MANAGEMENT

E. Chronic Care Improvement Program (CCIP)

- 2) The CCIP Year 1 Intervention Plan section
 - b. The CCIP Overview describes the 3-year CCIP focus, including the targeted chronic condition, population details, CCIP aim including the data source(s), and intervention strategy.
 - c. The CCIP Year 1 Intervention Plan section includes CCIP criteria for identifying the focus population, the goal, planned interventions, and measurement methodology including applicable data sources.
2. Subsequent Submissions – Due semi-annually following the first (1st) submission of that year. Refer to the table below for submission details.
 - a. The Progress Update describes the Delegate’s progress in implementing the CCIP, including systematic and ongoing follow-up.
 - 1) First (1st) semi-annual submissions to include:
 - Reporting CCIP intervention results, findings, and lessons learned, covering the Study/Act sections of the PDSA.
 - The Delegate’s Plan for the next CCIP cycle year.
 - 2) Second (2nd) semi-annual submissions to include:
 - Reporting the progress and status of the CCIP Plan, including timelines, barriers, and planned next steps.

CCIP Year	Submission*	Submission Component Due: (CCIP Cycle & PDSA Focus)
Year 1	1 st Semi-Annual	<i>CCIP Program Launch –</i> <ul style="list-style-type: none"> • CCIP Overview • Cycle 1 – Plan
	2 nd Semi-Annual	<ul style="list-style-type: none"> • Cycle 1 – Do
Year 2	1 st Semi-Annual	<ul style="list-style-type: none"> • Cycle 1– Study, Adjust/Act/Abandon • Cycle – 2 Plan
	2 nd Semi-Annual	<ul style="list-style-type: none"> • Cycle 2 – Do
Year 3	1 st Semi-Annual	<ul style="list-style-type: none"> • Cycle 2 – Study, Adjust/Act /Abandon • Cycle 3 – Plan
	2 nd Semi-Annual	<ul style="list-style-type: none"> • Cycle 3 – Do
Year 3 Final Closeout/ Launch New CCIP	1 st Semi-Annual	<i>CCIP Program Close-Out –</i> <ul style="list-style-type: none"> • Cycle 3 – Study, Act • CCIP Close Out <i>Launch NEW CCIP</i> <ul style="list-style-type: none"> • <i>Begin new CCIP Document</i>

13. QUALITY MANAGEMENT

E. Chronic Care Improvement Program (CCIP)

See Attachment, “Chronic Care Improvement Program (CCIP) Planning & Reporting Document” in Section 13 for submission due dates & documentation requirements.

F. IEHP and its IPAs must make information on the status and results of ongoing CCIP projects available to IEHP at any time, upon request.⁸

G. CMS Annual Reporting:

1. IEHP utilizes the Health Plan Management System (HPMS) to report the status of their CCIP to CMS by December 31st, annually. Submissions include an attestation by IEHP regarding its compliance with the ongoing CCIP requirement.

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Chief Approval: <i>Signature on File</i>	Original Effective Date:	January 1, 2023
Chief Title: Chief Medical Officer	Revision Date:	

⁸42 CFR § 422.152(f)(1)(i)(iii)

13. COORDINATION OF CARE

Attachments

<u>DESCRIPTION</u>	<u>POLICY CROSS REFERENCE</u>
Provider Preventable Conditions	13D
Chronic Care Improvement Program (CCIP) Planning & Reporting Document	13E

<p>Category 1 – Health Care-Acquired Conditions (For Any Inpatient Hospital Settings in Medicaid)</p>	<ul style="list-style-type: none"> • Any unintended foreign object retained after surgery • A clinically significant air embolism • An incident of blood incompatibility • A Stage III or IV pressure ulcer that developed during the patient’s stay in the hospital • A significant fall or trauma that resulted in fracture, dislocation, intracranial injury, crushing injury, burn, or electric shock • A catheter-associated urinary tract infection (UTI) • Vascular catheter-associated infection • Any of the following manifestations of poor glycemic control: diabetic ketoacidosis; nonketotic hyperosmolar coma; hypoglycemic coma; secondary diabetes with ketoacidosis; or secondary diabetes with hyperosmolarity • A surgical site infection following: <ul style="list-style-type: none"> ○ Coronary artery bypass graft (CABG) - mediastinitis ○ Bariatric surgery; including laparoscopic gastric bypass, gastroenterostomy, laparoscopic gastric restrictive surgery ○ Orthopedic procedures; including spine, neck, shoulder, elbow ○ Cardiac implantable electronic device (CIED) procedures • Deep vein thrombosis (DVT)/pulmonary embolism (PE) following total knee replacement with pediatric and obstetric exceptions • Iatrogenic pneumothorax with venous catheterization • A vascular catheter-associated infection
<p>Category 2 – Other Provider Preventable Conditions (For Any Health Care Setting)</p>	<ul style="list-style-type: none"> • Wrong surgical or other invasive procedure performed on a

	<p>patient</p> <ul style="list-style-type: none">• Surgical or other invasive procedure performed on the wrong body part• Surgical or other invasive procedure performed on the wrong patient
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Chronic Care Improvement Program (CCIP) Planning & Reporting Document

[IPA Name]

CCIP Reporting Period: 01/01/23 – 12/31/25

Cycle 1: 01/01/23 – 12/31/23

Cycle 2: 01/01/24 – 12/31/24

Cycle 3: 01/01/25 – 12/31/25

Final Submission: 03/15/26

Note: Do not include Member PHI in your summaries.

Table of Contents

PROGRAM YEAR 1:

Year 1, Cycle 1 – CCIP Overview & “Plan” –

Due to IEHP by: 03/15/2023 (1st Submission)

3-Year CCIP Overview:

Plan of CCIP Cycle 1 Intervention:

FOR IEHP INTERNAL USE ONLY – 1st Submission (CCIP Overview and Cycle 1 Plan)

Year 1, Cycle 1 – CCIP “DO” –

Due to IEHP by: 09/15/2023 (2nd Submission)

Progress Update of CCIP Cycle 1 Action:

FOR IEHP INTERNAL USE ONLY – 2nd Submission (Progress Update).

PROGRAM YEAR 2:

Year 2, Cycle 1 – CCIP “Study/Act” & Cycle 2 – CCIP “Plan” -

Due to IEHP by: 03/15/2024 (3rd Submission)

Analysis of CCIP Cycle 1 Intervention:

Plan of CCIP Cycle 2 Intervention:

FOR IEHP INTERNAL USE ONLY – 3rd Submission (Progress Update).

Year 2, Cycle 2 – CCIP “DO” –

Due to IEHP by: 09/15/2024 (4th Submission)

Progress Update of CCIP Cycle 2 Action:

FOR IEHP INTERNAL USE ONLY – 4th Submission (Progress Update).

PROGRAM YEAR 3:

Year 3: Cycle 2 – CCIP “Study/Act” & Cycle 3 – CCIP “Plan” -

Due to IEHP by: 03/15/2025 (5th Submission)

Analysis of CCIP Cycle 2 Intervention:

Plan of CCIP Cycle 3 Intervention:

FOR IEHP INTERNAL USE ONLY – 5th Submission (Progress Update).

Year 3, Cycle 3 – CCIP “DO”-

Due to IEHP by: 09/15/2025 (6th Submission)

Progress Update of CCIP Cycle 3 Action:

FOR IEHP INTERNAL USE ONLY – 6th Submission (Progress Update).



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Chronic Care Improvement Program (CCIP) Planning & Reporting Document

PROGRAM CLOSE:

Year 3 Wrap Up, Cycle 3 – CCIP “Study/Act” & CCIP Summary

Due to IEHP by: 03/15/2026 (7th Submission)

Analysis of CCIP Cycle 3 Intervention:

CCIP Close-Out – Summary of 3-Year CCIP Plan:

FOR IEHP INTERNAL USE ONLY – 7th Submission (Final CCIP Update & Close-Out).

APPENDIX

CCIP Submission Dates

[IPA Name]

[CCIP Title]

CCIP Reporting Period: 01/01/2023 – 12/31/2025

PROGRAM YEAR 1:

Year 1, Cycle 1 – CCIP Overview & “Plan” –

Due to IEHP by: 03/15/2023 (1st Submission)

3-Year CCIP Overview:

CCIP Overview	
Line of Business:	Medicare
Targeted Chronic Condition & Focus:	<p>Select ONE (1) focus opportunity from the options listed below:</p> <p>Diabetes:</p> <ul style="list-style-type: none"> <input type="checkbox"/> Diabetes Care – Eye Exam <input type="checkbox"/> Diabetes Care – Kidney Disease Monitoring <input type="checkbox"/> Diabetes Care – Blood Sugar Controlled (HbA1c >9%) <input type="checkbox"/> Statin Therapy for Patients with Diabetes <p>Cardiovascular Disease:</p> <ul style="list-style-type: none"> <input type="checkbox"/> Statin Therapy for Patients with Cardiovascular Disease <input type="checkbox"/> Controlling High Blood Pressure
Average IEHP D-SNP Population Size:	[12-month average]
CCIP Aim (Outcome Measure):	
[Aim to be written in “S.M.A.R.T” format: Specific, Measurable, Attainable, Relevant/Realistic, Timely]	
Baseline:	Target:
[N:D = Rate]	[Rate]
Data Source(s) to be Used in Evaluation of CCIP Performance	
[Refer to the CCIP Reference Guide for sample sources]	

Plan of CCIP Cycle 1 Intervention:

Intervention Details	
Intervention Name:	

[IPA Name]: [CCIP Title] – 01/01/23 – 12/31/25

Reminder: Do not include Member PHI in your summaries.



Chronic Care Improvement Program (CCIP) Planning & Reporting Document

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Planned Strategy:	[Refer to the CCIP Reference Guide for eligible options]
Intervention Description:	[Describe the current improvement opportunity you are looking to address through this intervention, including the potential impact to Members and/or Providers.]
Testing Period:	[MM/DD/YY – MM/DD/YY]
Measurement Methodology (Process Measure):	
[Describe how will you measure the success of this intervention. What tool(s)/report(s) will be used, how often it will be assessed, and with whom will the results be shared?]	
Reporting Frequency:	[Requirement: Measurements should be monitored monthly, at minimum.]
Description of Numerator:	[Describe the data to be measured at the numerator level]
Description of Denominator:	[Describe the data to be measured at the denominator level]
Baseline	Target
[N:D = Rate]	[Rate]
Intervention Process:	
1. [List the process steps of your intervention.]	

FOR IEHP INTERNAL USE ONLY – 1st Submission (CCIP Overview and Cycle 1 Plan)

Initial Plan Submission – Due 03/15/23			
CCIP Received by IEHP:			
Received Date:		By (i#):	
Quality Review:			
Reviewed Date:		P4P Quality Measure:	
		Timeliness:	<input type="checkbox"/> Met <input type="checkbox"/> Not Met
By (i#):		Completeness:	<input type="checkbox"/> Met <input type="checkbox"/> Not Met
Resubmission Required?	<input type="checkbox"/> No <input type="checkbox"/> Yes, due back to IEHP by: _____		
Notes:			

[IPA Name]: [CCIP Title] – 01/01/23 – 12/31/25

Reminder: Do not include Member PHI in your summaries.



Chronic Care Improvement Program (CCIP) Planning & Reporting Document

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Year 1, Cycle 1 – CCIP “DO” –

Due to IEHP by: 09/15/2023 (2nd Submission)

Progress Update of CCIP Cycle 1 Action:

Intervention Details	
Intervention Status:	<input type="checkbox"/> On Track – <i>progressing as scheduled</i> <input type="checkbox"/> Off Track – <i>progress is delayed/off schedule or has not begun</i>
Summary of Current Status:	[Describe the status of your current intervention. What has been done, results seen, and whether this intervention is progressing as planned?]
Barriers:	[Describe any barriers encountered and your mitigation strategies.]
Lessons Learned & Best Practices:	[Describe the lessons learned as you have begun executing your intervention. Provide any best practices you have adopted.]
Next Steps:	
[Describe the next steps to your intervention, including anticipated timeframes.]	

FOR IEHP INTERNAL USE ONLY – 2nd Submission (Progress Update).

Progress Update Submission – Due 09/15/23			
CCIP Received by IEHP:			
Received Date:		By (i#):	
Quality Review:			
Reviewed Date:		P4P Quality Measure:	
		Timeliness:	<input type="checkbox"/> Met <input type="checkbox"/> Not Met
By (i#):		Completeness:	<input type="checkbox"/> Met <input type="checkbox"/> Not Met
Resubmission Required?	<input type="checkbox"/> No <input type="checkbox"/> Yes, due back to IEHP by: _____		
Notes:			

[IPA Name]: [CCIP Title] – 01/01/23 – 12/31/25

Reminder: Do not include Member PHI in your summaries.



PROGRAM YEAR 2:

Year 2, Cycle 1 – CCIP “Study/Act” & Cycle 2 – CCIP “Plan” -

Due to IEHP by: 03/15/2024 (3rd Submission)

Analysis of CCIP Cycle 1 Intervention:

Intervention Details	
Intervention Results:	<input type="checkbox"/> Met – <i>Target goal was achieved</i> <input type="checkbox"/> Not Met – <i>Target goal was not achieved</i>
Intervention Results:	
Baseline (from above):	Target (from above):
[N:D = Rate]	[Rate]
Actual:	[N:D = Rate]
Results and Findings:	[Summarize the results and findings of the intervention. Describe using qualitative and quantitative data.]
Barriers:	[Describe any new barriers encountered and your mitigation strategies.]
Lessons Learned & Best Practices:	[Describe the lessons learned as you completed your intervention. Provide any new best practices you have adopted.]
Next Steps for this Intervention:	
Next Steps leading into Year 2, Cycle 2:	<input type="checkbox"/> Adopt – <i>Intervention is ready for integration.</i> <input type="checkbox"/> Adjust – <i>Intervention needs modifications.</i> <input type="checkbox"/> Abandon – <i>Intervention to conclude with no further action.</i> <input type="checkbox"/> Continue – <i>Would like to proceed with further testing.</i>

Plan of CCIP Cycle 2 Intervention:

Intervention Details	
Intervention Name:	
Planned Strategy:	[Refer to the CCIP Reference Guide for eligible options]
Intervention Description:	[Describe the current improvement opportunity you are looking to address through this intervention, including the potential impact to Members and/or Providers.]
Testing Period:	[MM/DD/YY – MM/DD/YY]
Measurement Methodology (Process Measure):	
[Describe how will you measure the success of this intervention. What tool(s)/report(s) will be used, how often it will be assessed, and with whom will the results be shared?]	
Reporting Frequency:	[Requirement: Measurements should be monitored monthly, at minimum.]
Description of Numerator:	[Describe the data to be measured at the numerator level]

[IPA Name]: [CCIP Title] – 01/01/23 – 12/31/25

Reminder: Do not include Member PHI in your summaries.



Chronic Care Improvement Program (CCIP) Planning & Reporting Document

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Description of Denominator:	[Describe the data to be measured at the denominator level]	
	Baseline	Target
	[N:D = Rate]	[Rate]
Intervention Process:		
1. [List the process steps of your intervention.]		

FOR IEHP INTERNAL USE ONLY – 3rd Submission (Progress Update).

Progress Update Submission – Due 03/15/24			
CCIP Received by IEHP:			
Received Date:		By (i#):	
Quality Review:			
Reviewed Date:		P4P Quality Measure:	
		Timeliness:	<input type="checkbox"/> Met <input type="checkbox"/> Not Met
By (i#):		Completeness:	<input type="checkbox"/> Met <input type="checkbox"/> Not Met
Resubmission Required?	<input type="checkbox"/> No <input type="checkbox"/> Yes, due back to IEHP by: _____		
Notes:			



Chronic Care Improvement Program (CCIP) Planning & Reporting Document

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Year 2, Cycle 2 – CCIP “DO” –

Due to IEHP by: 09/15/2024 (4th Submission)

Progress Update of CCIP Cycle 2 Action:

Intervention Details	
Intervention Status:	<input type="checkbox"/> On Track – <i>progressing as scheduled</i> <input type="checkbox"/> Off Track – <i>progress is delayed/off schedule or has not begun</i>
Summary of Current Status:	[Describe the status of your current intervention. What has been done, results seen, and whether this intervention is progressing as planned?]
Barriers:	[Describe any barriers encountered and your mitigation strategies.]
Lessons Learned & Best Practices:	[Describe the lessons learned as you have begun executing your intervention. Provide any best practices you have adopted.]
Next Steps:	
[Describe the next steps to your intervention, including anticipated timeframes.]	

FOR IEHP INTERNAL USE ONLY – 4th Submission (Progress Update).

Progress Update Submission – Due 09/15/24			
CCIP Received by IEHP:			
Received Date:		By (i#):	
Quality Review:			
Reviewed Date:		P4P Quality Measure:	
		Timeliness:	<input type="checkbox"/> Met <input type="checkbox"/> Not Met
By (i#):		Completeness:	<input type="checkbox"/> Met <input type="checkbox"/> Not Met
Resubmission Required?	<input type="checkbox"/> No <input type="checkbox"/> Yes, due back to IEHP by: _____		
Notes:			

[IPA Name]: [CCIP Title] – 01/01/23 – 12/31/25

Reminder: Do not include Member PHI in your summaries.



PROGRAM YEAR 3:

Year 3: Cycle 2 – CCIP “Study/Act” & Cycle 3 – CCIP “Plan” -

Due to IEHP by: 03/15/2025 (5th Submission)

Analysis of CCIP Cycle 2 Intervention:

Intervention Details		
Intervention Results:	<input type="checkbox"/> Met – <i>Target goal was achieved</i> <input type="checkbox"/> Not Met – <i>Target goal was not achieved</i>	
Intervention Results:		
Baseline (from above):	Target (from above):	Actual:
[N:D = Rate]	[Rate]	[N:D = Rate]
Results and Findings:	[Summarize the results and findings of the intervention. Describe using qualitative and quantitative data.]	
Barriers:	[Describe any new barriers encountered and your mitigation strategies.]	
Lessons Learned & Best Practices:	[Describe the lessons learned as you completed your intervention. Provide any new best practices you have adopted.]	
Next Steps for this Intervention:		
Next Steps leading into Year 2, Cycle 2:	<input type="checkbox"/> Adopt – <i>Intervention is ready for integration.</i> <input type="checkbox"/> Adjust – <i>Intervention needs modifications.</i> <input type="checkbox"/> Abandon – <i>Intervention to conclude with no further action.</i> <input type="checkbox"/> Continue – <i>Would like to proceed with further testing.</i>	

Plan of CCIP Cycle 3 Intervention:

Intervention Details	
Intervention Name:	
Planned Strategy:	[Refer to the CCIP Reference Guide for eligible options]
Intervention Description:	[Describe the current improvement opportunity you are looking to address through this intervention, including the potential impact to Members and/or Providers.]
Testing Period:	[MM/DD/YY – MM/DD/YY]
Measurement Methodology (Process Measure):	
[Describe how will you measure the success of this intervention. What tool(s)/report(s) will be used, how often it will be assessed, and with whom will the results be shared?]	
Reporting Frequency:	[Requirement: Measurements should be monitored monthly, at minimum.]
Description of Numerator:	[Describe the data to be measured at the numerator level]

[IPA Name]: [CCIP Title] – 01/01/23 – 12/31/25

Reminder: Do not include Member PHI in your summaries.



Chronic Care Improvement Program (CCIP) Planning & Reporting Document

DualChoice

Description of Denominator:	[Describe the data to be measured at the denominator level]	
	Baseline	Target
	[N:D = Rate]	[Rate]
Intervention Process:		
1. [List the process steps of your intervention.]		

FOR IEHP INTERNAL USE ONLY – 5th Submission (Progress Update).

Progress Update Submission – Due 03/15/25			
CCIP Received by IEHP:			
Received Date:		By (i#):	
Quality Review:			
Reviewed Date:		P4P Quality Measure:	
By (i#):		Timeliness:	<input type="checkbox"/> Met <input type="checkbox"/> Not Met
Resubmission Required?	<input type="checkbox"/> No <input type="checkbox"/> Yes, due back to IEHP by: _____		
Completeness:			<input type="checkbox"/> Met <input type="checkbox"/> Not Met
Notes:			

[IPA Name]: [CCIP Title] – 01/01/23 – 12/31/25

Reminder: Do not include Member PHI in your summaries.



Chronic Care Improvement Program (CCIP) Planning & Reporting Document

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Year 3, Cycle 3 – CCIP “DO”-

Due to IEHP by: 09/15/2025 (6th Submission)

Progress Update of CCIP Cycle 3 Action:

Intervention Details	
Intervention Status:	<input type="checkbox"/> On Track – <i>progressing as scheduled</i> <input type="checkbox"/> Off Track – <i>progress is delayed/off schedule or has not begun</i>
Summary of Current Status:	[Describe the status of your current intervention. What has been done, results seen, and whether this intervention is progressing as planned?]
Barriers:	[Describe any barriers encountered and your mitigation strategies.]
Lessons Learned & Best Practices:	[Describe the lessons learned as you have begun executing your intervention. Provide any best practices you have adopted.]
Next Steps:	
[Describe the next steps to your intervention, including anticipated timeframes.]	

FOR IEHP INTERNAL USE ONLY – 6th Submission (Progress Update).

Progress Update Submission – Due 09/15/25			
CCIP Received by IEHP:			
Received Date:		By (i#):	
Quality Review:			
Reviewed Date:		P4P Quality Measure:	
		Timeliness:	<input type="checkbox"/> Met <input type="checkbox"/> Not Met
By (i#):		Completeness:	<input type="checkbox"/> Met <input type="checkbox"/> Not Met
Resubmission Required?	<input type="checkbox"/> No <input type="checkbox"/> Yes, due back to IEHP by: _____		
Notes:			

[IPA Name]: [CCIP Title] – 01/01/23 – 12/31/25

Reminder: Do not include Member PHI in your summaries.



PROGRAM CLOSE:

Year 3 Wrap Up, Cycle 3 – CCIP “Study/Act” & CCIP Summary

Due to IEHP by: 03/15/2026 (7th Submission)

Analysis of CCIP Cycle 3 Intervention:

Intervention Details	
Intervention Results:	<input type="checkbox"/> Met – <i>Target goal was achieved</i> <input type="checkbox"/> Not Met – <i>Target goal was not achieved</i>
Intervention Results:	
Baseline (from above):	Target (from above):
[N:D = Rate]	[Rate]
	Actual:
	[N:D = Rate]
Results and Findings:	[Summarize the results and findings of the intervention. Describe using qualitative and quantitative data.]
Barriers:	[Describe any new barriers encountered and your mitigation strategies.]
Lessons Learned & Best Practices:	[Describe the lessons learned as you completed your intervention. Provide any new best practices you have adopted.]
Final Steps for this Intervention:	
Final Steps:	<input type="checkbox"/> Adopt – <i>Intervention is ready for integration.</i> <input type="checkbox"/> Adjust – <i>Intervention needs modifications.</i> <input type="checkbox"/> Abandon – <i>Intervention to conclude with no further action.</i> <input type="checkbox"/> Continue – <i>Would like to proceed with further testing.</i>

CCIP Close-Out – Summary of 3-Year CCIP Plan:

CCIP Details:			
CCIP Results:	<input type="checkbox"/> Met – <i>Aim was achieved</i> <input type="checkbox"/> Not Met – <i>Aim was not achieved</i>		
CCIP SMART Aim Attainment:			
CCIP SMART Aim	Baseline	Target	Actual
	[N:D= %]	[%]	[N:D= %]
Results of the Intervention:			
Results and Findings:	[Summarize the results and findings of the overall CCIP plan. Describe using qualitative and quantitative data.]		
Barriers:	[Describe any major barriers to your program.]		
Lessons Learned & Best Practices:	[Describe the lessons learned throughout this CCIP Process, including what may be done differently in the future. Include the best practices you have adopted from this CCIP experience.]		

[IPA Name]: [CCIP Title] – 01/01/23 – 12/31/25

Reminder: Do not include Member PHI in your summaries.



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DualChoice

Closing Remarks:
[Free text: Include any closing remarks or insights related to this 3-year CCIP experience, including insights.]

FOR IEHP INTERNAL USE ONLY – 7th Submission (Final CCIP Update & Close-Out).

Final Submission – Due 03/15/26			
CCIP Received by IEHP:			
Received Date:		By (i#):	
Quality Review:			
Reviewed Date:		P4P Quality Measure:	
		Timeliness:	<input type="checkbox"/> Met <input type="checkbox"/> Not Met
By (i#):		Completeness:	<input type="checkbox"/> Met <input type="checkbox"/> Not Met
Resubmission Required?	<input type="checkbox"/> No <input type="checkbox"/> Yes, due back to IEHP by: _____		
Notes:			

[IPA Name]: [CCIP Title] – 01/01/23 – 12/31/25

Reminder: Do not include Member PHI in your summaries.



APPENDIX

CCIP Submission Dates

CCIP progress is to be submitted to IEHP semi-annually, over the course of three (3) years, with a final reflection in the 4th year.

Refer to the table below for submission details, including reflection periods, due dates and submission components.

CCIP Year	Submission / Reflection Period	Submission Due Date:	Submission Component Due: (CCIP Cycle & PDSA Focus)
Year 1	1 st Semi-Annual 01/01/23 – 02/28/23	03/15/23	1 st Submission: CCIP Program Launch – <ul style="list-style-type: none"> CCIP Overview Cycle 1 – Plan
	2 nd Semi-Annual 04/01/23 – 08/30/23	09/15/23	2 nd Submission: Progress Update – <ul style="list-style-type: none"> Cycle 1 – Do
Year 2	1 st Semi-Annual 09/01/23 – 02/29/24	03/15/24	3 rd Submission: Progress Update – <ul style="list-style-type: none"> Cycle 1– Study, Adjust/Act/Abandon Cycle – 2 Plan
	2 nd Semi-Annual 03/01/24 – 08/30/24	09/15/24	4 th Submission: Progress Update – <ul style="list-style-type: none"> Cycle 2 – Do
Year 3	1 st Semi-Annual 09/01/24 – 02/28/25	03/15/25	5 th Submission: Progress Update – <ul style="list-style-type: none"> Cycle 2 – Study, Adjust/Act /Abandon Cycle 3 – Plan
	2 nd Semi-Annual 03/01/25 – 08/30/25	09/15/25	6 th Submission: Progress Update – <ul style="list-style-type: none"> Cycle 3 – Do
Year 3 Final Closeout/ Launch New CCIP	1 st Semi-Annual 09/01/25 – 12/31/25 (CCIP Close Out) 01/01/26 – 02/28/26 (NEW CCIP)	03/15/26	7 th Submission: CCIP Program Close-Out – <ul style="list-style-type: none"> Cycle 3 – Study, Act CCIP Close Out Launch NEW CCIP <ul style="list-style-type: none"> Begin new CCIP Document

[IPA Name]: [CCIP Title] – 01/01/23 – 12/31/25

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