Health Information Exchange in California

Gaps and Opportunities
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Introduction

The exchange of information is essential to a well-functioning health and human service ecosystem and lays the foundation for the coordinated delivery of care and services to support health and well-being for individuals and communities. The sharing, collection, management, and use of health and human service data helps ensure that the individuals and communities that are at the center of the ecosystem can benefit from informed decisions based on complete information gathered across organizations and sectors.

In recent decades, exchange of health and human service information has grown. Federal and state government initiatives such as the Health Information Technology for Economic and Clinical Health Act of 2009 (HITECH Act) released billions of dollars in grants, loans, and incentives to accelerate adoption of technology and encourage information exchange.¹ Today, health care providers are largely using electronic systems to document health information, government and industry organizations have driven forward data exchange standards, and health information organizations and other data exchange intermediaries have emerged to support data sharing between entities and sectors.

Yet challenges to robust and effective health and human service data exchange persist. Not all health and human service providers in California today are able to access and share medical, behavioral, and social service histories.² Where such data exchange is occurring, it is often incomplete and confined to a subset of traditional care data, shared through varied mechanisms that do not support a coordinated approach for sharing data with all relevant parties, and limited by diverging interpretations of data sharing laws and regulations.

This document describes a set of gaps hampering robust and effective data sharing in California as well as associated opportunities that can address these gaps to further data exchange in the state. These gaps and opportunities describe input, recommendations, and other considerations provided by members of the Data Exchange Framework (DxF) Stakeholder Advisory Group and members of the public between August 2021 and June 2022. The Stakeholder Advisory Group conducted its business through discussion and consensus building, identifying and documenting key considerations of various DxF recommendations for advancement to the Secretary of the California Health and Human Services Agency (CalHHS). In the event that

consensus could not be reached, options, pros, and cons were advanced for consideration by the Secretary.

Identified gaps and opportunities fell into six categories:

- Technical Infrastructure and Health Information Technology (HIT) Capacity
- Data Exchange Standards
- Provider Information and Person Identity Management
- Individual Data Access
- Data Exchange Law, Regulations, and Policy
- Health and Human Service Information Exchange Financing

Together, the gaps and opportunities described in this paper represent a set of priority policy recommendations that California’s policymakers and stakeholders can use as a roadmap to advance robust and effective data exchange in the state. This document proposes recommendations that can serve as an important next step in improving meaningful data exchange and supporting health and well-being for all Californians.
Executive Summary

The following tables provide summaries of each of the gaps and opportunities included in this document. Further detail on each gap and opportunity is included in its corresponding section.

<table>
<thead>
<tr>
<th>A. Technical Infrastructure and Health Information Technology (HIT) Capacity</th>
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<tr>
<td><strong>Gap #1: Electronic Health Record (EHR) Adoption.</strong></td>
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<tr>
<td>Many health and human service organizations in California</td>
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<td>do not have the resources to invest in foundational digital</td>
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<td>record systems, which are required for the electronic</td>
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<td>collection and exchange of health information to support</td>
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<td>effective health service delivery. Incomplete adoption</td>
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<td>leaves critical individual/client data siloed, limiting cross-</td>
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<td>organization and cross-sector care coordination.</td>
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<td><strong>Opportunity #1.1: EHR Incentive Program.</strong> The state</td>
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<td>should establish an EHR incentive program to encourage</td>
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<td>adoption of certified EHR technology (CEHRT) and clinical</td>
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<td>documentation technologies among health care</td>
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<td>organizations that do not have the required technological</td>
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<td>capacity to support the collection, exchange, and use of</td>
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<tr>
<td>electronic health information in accordance with the</td>
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<td>California Health and Human Services Agency’s (CalHHS’s) Data</td>
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<td>Exchange Framework (DxF) requirements.</td>
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</table>
**Gap #2: Data Exchange Capacity at Many Health Care and Human Service Organizations.** Effective health and human service information exchange requires investment in interoperable technology capable of supporting data exchange (see Gap #A1) as well as the ability to connect and share information with other participants. However, many health and human service organizations confront technical, operational, and financial barriers to making those connections.

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<tr>
<th>Opportunity #2.1: Data Exchange Intermediary Onboarding and Technical Assistance Program for Health and Human Service Organizations. The state should establish a data exchange onboarding and technical assistance program that provides funding, incentives, and technical assistance to help health and human service organizations connect to a qualifying data exchange intermediary.</th>
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<tr>
<td>Opportunity #2.2: Data Exchange Intermediary Qualification Process. The state should establish policy that leverages national programs that define a qualification approach for data exchange intermediaries and should further specify additional state data sharing requirements pursuant to AB133 that should be incorporated into the DxF Data Sharing Agreement (DSA) and Policies and Procedures (P&amp;Ps).</td>
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**Gap #3: Event Notifications.** Event notifications – real-time, automated messages that can alert an individual’s primary care practitioner, care manager, or care team of changes to the individual’s condition (e.g., admission to, discharge from, or transfer to [ADT] a health care facility) – are a primary example of how health and human service information exchange can be used to support care coordination and service delivery, though their utilization is often limited to a narrow set of participants and circumstances (e.g., alerting of transitions from acute care facilities).

| Opportunity #3.1: Expansions to Federal Event Notification Requirements. The state should develop policy and contracting requirements that extend the scope of federal event notification requirements included in the Centers for Medicare & Medicaid Services (CMS) Interoperability and Patient Access final rule to include additional health and human service organizations and notification types. |
**Gap #4: Intra- and Inter-Sector Data Exchange Capabilities.** California’s public health and human service agencies are stewards of valuable information that they need to manage programs and services they deliver to clients, individuals, and families. Most have mixed capabilities to electronically exchange timely and usable information with other health and human service organizations that could benefit from accessing it to provide a more complete picture and help them deliver a richer complement of coordinated services.

**Opportunity #4.1: Upgrades to California County HIT Infrastructure.** The state should leverage and expand federally funded programs to upgrade state and local public health information technology infrastructure and to provide a glidepath for county health, public health, and human service entities to participate in information exchange.

**Opportunity #4.2: Public Agency Data Exchange Policy and Contracting Requirements.** The state should, through policy, procurement processes, and contract amendments, contractually obligate vendors to share information with health and human service organizations to advance goals envisioned by AB133.
### B. Data Exchange Standards

<table>
<thead>
<tr>
<th>Gap #1: Demographic and Human Service Data Exchange Standards and Capacity. The standardized collection, curation, and use of demographic and social determinants of health data in California remain uneven and developing.</th>
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<tbody>
<tr>
<td>Opportunity #1.1: Demographic Data Collection and Use Standards, Requirements, and Incentives. The state should establish demographic data collection and sharing requirements and incentives through public and private payers to encourage data use among health and human service organizations in California.</td>
</tr>
<tr>
<td>Opportunity #1.2: Social Determinants of Health (SDOH) Data Collection and Use Standards, Requirements, and Incentives. The state should establish SDOH data collection and sharing requirements and incentives through public and private payers to encourage data use among health and human service organizations in California.</td>
</tr>
<tr>
<td>Opportunity #1.3: Cross-State Agency Data Sharing Enhancements. The state should strengthen data exchange relationships between state agencies, supporting the exchange of program enrollment and relevant demographic and SDOH information to support a coordinated approach to health care and human service delivery across disparate programs and settings.</td>
</tr>
<tr>
<td>Opportunity #1.4: Recommendations to the Federal Government to Improve Demographic and SDOH Data Collection Nationally. The state should develop and advance recommendations to the federal government to improve demographic and SDOH data exchange standards nationally.</td>
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## C. Provider Information and Person Identity Management

<table>
<thead>
<tr>
<th>Gap #1: Robust Provider Information. Complete provider information (e.g., name, practice locations, organizational affiliates, and the available means to exchange health information such as Direct addresses and endpoints) is not always available or accessible to all health and human service organizations across California, creating barriers to effective information exchange and service delivery.</th>
<th>Opportunity #1.1: Provider Directory and Reporting. The state should support the establishment of a statewide provider directory and should require signatories of the DxF DSA to contribute data to encourage provider-to-provider communication and information exchange.</th>
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<tr>
<td>Gap #2: Person Identity Management. No robust or systematic coordination of digital identities, person resolution, or data linking exists across organizational boundaries in California, limiting the efficacy of cross-organizational data exchange. As a result, organizations often fail to locate existing health records for individuals they serve that might exist at other organizations to support care coordination and management, because the organizations’ health information systems fail to agree on a single personal “identity.”</td>
<td>Opportunity #2.1: Strategy for Digital Identities. The state should adopt the Strategy for Digital Identities [final link to be inserted] called for in AB133 as a component of the DxF.</td>
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## D. Individual Data Access

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<thead>
<tr>
<th>Gap #1: Individual Data Access</th>
<th>Opportunity #1.1: Policies to Ensure That Individuals Have Meaningful Access to Their Longitudinal Health Information</th>
<th>Opportunity #1.2: Policies to Strengthen Understanding and Trust for Individuals Sharing Data</th>
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<tr>
<td>Individuals consistently face challenges in accessing and contributing to their health and human service records – directly or through selected third parties – in a manner that is convenient, timely, and compliant with federal access requirements.</td>
<td>The state should adopt and expand policies to ensure that individuals have meaningful access to their longitudinal health information across all health care organizations that are required to execute the DxF DSA, allowing all Californians to be meaningfully engaged in their care and make informed health care decisions.</td>
<td>The state should assess, strengthen, and adopt policies that will support individuals in understanding how their health and human service information may be used and will help establish trust in the systems in place that govern its use.</td>
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E. Data Exchange Law, Regulations, and Policy

| Gap #1: Data Exchange Law, Regulations, and Policy. Numerous federal and state laws, regulations, and policies that govern the exchange of physical, behavioral health, and human service data create real or perceived barriers to sharing information that is necessary to inform whole person care and population health goals. |
| Opportunity #1.1: “Universal” Release-of-Information Authorization Form. The state should support the adoption and use of a “universal” release-of-information authorization form to enable standardized data exchange in support of state priority use cases and the goals of AB133. |
| Opportunity #1.2: Consent Management Service: The state should consider support for a consent management service that would allow individuals to electronically manage – and health and human service organizations to access – consents to disclose and share certain types of health and human service information. |

F. Health and Human Service Information Exchange Financing

| Gap #1: Financing to Support Health and Human Service Information Exchange. Many barriers to data exchange are rooted in challenges for health and human service organizations in accessing financing to make needed health information technology investments. Funding is needed to establish an environment in California that is supportive of health and human service data exchange and to provide direct support for the organizations participating in exchange, particularly those that are under-resourced. |
| Opportunity #1.1: Leveraging Governmental, Private, and Philanthropic Sources of Funding. The state should identify and pursue federal, state, private, and philanthropic funding opportunities to finance data exchange priorities consistent with the vision and goals of the DxF and AB133. |
A. Technical Infrastructure and Health Information Technology (HIT) Capacity

Gap #1: Electronic Health Record (EHR) Adoption

**Relevant AB133 Provisions**
Identify gaps, and propose solutions to gaps, in the life cycle of health information, including gaps in:

- [the] storage, maintenance, and management of health information. [§130290(c)(3)(B)(iii)]
- Linking, sharing, exchanging, and providing access to health information. [§130290(c)(3)(B)(iv)]

Many health and human service organizations in California do not have the resources to invest in foundational digital record systems, which are required for the electronic collection and exchange of health information to support effective health service delivery. Incomplete adoption leaves critical individual/client data siloed, limiting cross-organization and cross-sector care coordination.

Electronic health record (EHR) adoption has grown significantly since the HITECH Act, with the vast majority of hospitals and physicians in California now having an EHR (83% of hospitals in 2019; 79% of physicians in 2017). However, EHR adoption continues to lag at smaller and less well-resourced providers, as well as among providers that did not previously have access to financing from HITECH and other federal and state modernization funding opportunities (e.g., behavioral health organizations, long-term care facilities, correctional facility health providers). For example, only 84% of psychiatric hospitals nationally possess an EHR compared to 96% of general acute care hospitals, according to data from 2019 to 2021.

Further, EHR adoption does not guarantee that implemented systems have the capability to effectively collect, store, curate, and exchange data in line with current national interoperability standards, as established by the Office of the National Coordinator for Health Information Technology (ONC), or that they are connected to networks capable of health information exchange (see Gap #A2). Viewing adoption of certified EHR technology (CEHRT) reveals only further disparities between providers that were previously eligible for federal and state funding opportunities and those that

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3 The EHR adoption rate among hospitals was calculated using responses to the American Hospital Association (AHA) Annual Survey’s IT Supplement, which changed response options in 2019. The change likely contributed to a fall in reported adoption rates (from 97% in 2018 to 83% in 2019).
5 Office of the National Coordinator for Health Information Technology, “Adoption of Electronic Health Records by Hospital Service Type 2019-2021,” Health IT Quick Stat #60, March 28, 2022.
were not: only 75% of psychiatric hospitals nationally have CEHRT, compared to 94% of general acute care hospitals.\(^6\)

**Opportunity #1.1: EHR Incentive Program**

**Summary:** The state should establish an EHR incentive program to encourage adoption of certified EHR technology and clinical documentation technologies among health care organizations that do not have the required technological capacity to support the collection, exchange, and use of electronic health information in accordance with DxF requirements.

The state should establish an EHR incentive program that would encourage health care organizations to invest in EHR and clinical documentation technology capable of collecting, exchanging, and using electronic health information pursuant to CalHHS DxF requirements. EHRs have been shown to help improve care by enhancing information access, reducing preventable medical errors, and providing clinical decision support. EHRs can also improve care delivery efficiency, with one 2019 meta-analysis finding 7.4% lower costs among hospitals with EHRs compared to hospitals without EHRs.\(^7\)

EHR incentive programs have proven effective in increasing EHR adoption: one 2017 study found that annual EHR adoption rates for hospitals that were eligible for HITECH funding were 7.9 percentage points higher in the period after HITECH implementation compared to hospitals that were ineligible.\(^8\)

Other states have similarly developed and implemented EHR incentive programs to close critical health information collection and exchange gaps in their health systems:

- **North Carolina’s** EHR Incentive Program for Behavioral Health/IDD Providers was launched in May 2018 by the North Carolina Department of Health and Human Services Office of Rural Health, Division of Medical Assistance, and the North Carolina Health Information Exchange Authority. The program provided behavioral health, mental health, and intellectual development and disability practices with funding to support the purchase of EHR technology and connection to NC HealthConnex, the state-designated health information exchange (HIE). State law requires that most providers of Medicaid and state-funded health care services, even those without an EHR, connect to the state’s HIE by January 1, 2023.\(^9\)

- **New Jersey’s** Substance Use Disorder (SUD) Promoting Interoperability Program, funded by the New Jersey Department of Health and Department of Human Services, was established to provide funding to qualifying SUD treatment

\(^6\) Ibid.


providers to support the deployment or upgrading of EHR technology. Enrollment in the program began in 2019 and was planned to extend through June 2023, with funding disbursed on a first-come, first-served basis.

The scope of California’s EHR incentive program should include health care organizations with a demonstrated financial need that are required by AB133 to execute the DxF Data Sharing Agreement (DSA). The program should prioritize support for organizations that were not eligible for HITECH funding, such as acute psychiatric hospitals and certain behavioral health providers, and should ensure that all funding is used to support the acquisition, enhancement, and implementation of technologies that can support California’s DxF DSA requirements. The state should provide guidance that specifies minimum EHR technology and service requirements to meet stipulations of the DxF and its DSA.

A state EHR incentive program would be most effective if paired with investments and requirements advanced by public and private payers. Public and private payers, including Covered California, Medi-Cal, California Public Employees’ Retirement System (CalPERS), and commercial health plans, should consider establishing or reinforcing value-based payment arrangements that incentivize or contractually require health provider adoption of technologies that support health information exchange.

The state, in coordination with public and private payers, should also elevate persistent and pervasive industry health information technology gaps to the relevant federal departments (e.g., the Centers for Medicare & Medicaid Services [CMS], ONC) and should advocate for broader federal funding to close gaps.

A state EHR incentive program would complement and reinforce significant system reforms and investments already underway in California – including Medi-Cal’s CalAIM, which provides information technology and exchange investments through its Incentive Payment Program, the Providing Access and Transforming Health (PATH) expenditure authority, and the Behavioral Health Quality Incentive Program. Coordinated state investments and oversight will be critical to building the statewide health information technology capacity needed to support effective health information exchange.

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10 E.g., CEHRT for covered entities as defined by the Health Insurance Portability and Accountability Act (HIPAA) or other clinical documentation technologies for organizations that are not covered by HIPAA.


Gap #2: Data Exchange Capacity at Many Health Care and Human Service Organizations

**Relevant AB133 Provisions**

Identify gaps, and propose solutions to gaps, in the life cycle of health information, including gaps in linking, sharing, exchanging, and providing access to health information.  [§130290(c)(3)(B)(iv)]

Effective health and human service information exchange requires investment in interoperable technology capable of supporting data exchange (see Gap #A1) as well as the ability to connect and share information with other participants. However, many health and human service organizations confront technical, operational, and financial barriers to making those connections.

In California, many health and human service organizations have yet to make the necessary connections to enable meaningful data exchange. Regional health information organizations (HIOs) exchange data with fewer than half of the state’s hospitals and do not have a presence in approximately 19 of the state’s 58 counties.\(^ {14,15}\) National networks also present challenges to connection, with each network varying in the types of entities it supports and often requiring organizations to leverage internal information technology resources, inhibiting participation of small or under-resourced providers.\(^ {16}\) Where connections to HIOs, national networks, or other data exchange intermediaries are possible, providers may not be connected due to:

- A lack of understanding of the value health information exchange can bring to their patient care, and potentially to their practices through payer and provider incentives that may be in place;
- Expected challenges with meaningfully integrating such information into their clinical and care management workflows;
- Technical challenges in reconfiguring their EHR systems to connect to a data exchange intermediary;
- Legal concerns – wanting to ensure that potential data access and exchange is in compliance with existing federal and state law as well as any existing contractual and data sharing requirements the entity may have; and
- The costs of addressing these concerns, as well as paying any fees associated with onboarding to a data exchange intermediary and ongoing participation.

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\(^ {16}\) Ibid.
Comprehensive health and human service data exchange – even among connected health and human service organizations – may still be limited by data exchange intermediaries’ ability to support the spectrum of data exchange envisioned by federal and state regulations, and the aspirations of AB133 and many health care payers and providers. Data exchange intermediaries – both national and regional networks – vary in their capacity to exchange behavioral health data that may otherwise be protected by federal law (e.g., 42 C.F.R. Part 2); exchange social determinants of health data, where federal standards have only been recently released (e.g., United States Core Data for Interoperability [USCDI] version 2\textsuperscript{17}) and participant adoption of these standards is limited; and offer broader connections to individuals’ information than their direct network allows.\textsuperscript{18,19}

Opportunity #2.1: Data Exchange Intermediary Onboarding and Technical Assistance Program for Health and Human Service Organizations

**Summary:** The state should establish a data exchange onboarding and technical assistance program that provides funding, incentives, and technical assistance to help health and human service organizations connect to a qualifying data exchange intermediary.

The state should support the establishment of a data exchange intermediary onboarding program to increase statewide participation in health information exchange among health and human service organizations.

The program should prioritize investments in under-resourced health and human service providers, particularly those serving high-need, low-income, and historically disadvantaged populations, seeking connection to “qualified” data exchange intermediaries (see Opportunity #2.2).

The state should establish a technical assistance “Center of Excellence” to provide support for health and human service providers’ initial onboarding, potentially through a statewide initiative funded in collaboration with philanthropic organizations as well as private and public payers. The onboarding program would also support payment to connect organizations and qualified data exchange intermediaries, but would not be used to pay ongoing costs associated with maintaining connection to a data exchange intermediary. The onboarding program may build upon program predecessors, including the California HIE Onboarding Program (Cal-HOP), a Department of Health Care Services (DHCS) health information exchange initiative that provided state and


federal funding to help Medi-Cal providers onboard to Qualified Health Information Organizations (QHIOs). Early research on the Cal-HOP program, which concluded in September 2021, found that 89% of ambulatory providers and 79% of hospitals met all three program milestones and that HIOs participating in Cal-HOP generally perceived the program to be successful in supporting some provider organizations to exchange data via an HIO. Cal-HOP, among other state health or human service onboarding programs, should be assessed in advance of program launch for lessons learned. Behavioral health organizations, for example, may require a specific type of onboarding support to address challenges or concerns with managing data sharing consent and authorizations related to 42 C.F.R. Part 2.

A state data exchange intermediary onboarding program to support information exchange would complement and reinforce significant system reforms and investments presently underway in California (e.g., Medi-Cal’s CalAIM Incentive Payment Program, PATH, and the Behavioral Health Quality Incentive Program), and would ideally be implemented in partnership with parallel private-sector investments in data exchange intermediary onboarding.

Opportunity #2.2: Data Exchange Intermediary Qualification Process

**Summary:** The state should establish policy that leverages national programs that define a qualification approach for data exchange intermediaries and should further specify additional state data sharing requirements pursuant to AB133 that should be incorporated into the DxF DSA and Policies and Procedures.

The state should establish a formal process for qualifying data exchange intermediaries that meet CalHHS DxF requirements. The qualification process should build on federal and state models, including the federal Trusted Exchange Framework and Common Agreement’s (TEFCA) concept of a Qualified Health Information Network (QHIN) as

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21 The goals of Cal-HOP were to (1) increase the number of Medi-Cal providers that can exchange patient data via a health information organization (HIO); (2) expand the data exchange capabilities of Medi-Cal providers already participating in HIOs; and (3) facilitate Medi-Cal providers’ access to the Controlled Substance Utilization Review and Evaluation System (CURES) prescription drug monitoring database. The program launched in 2019, and project activity concluded effective September 30, 2021.
26 A QHIN is a network designated as having met specified criteria described in TEFCA. QHIN criteria include the ability to exchange specified required information and perform required functions, as well as possession of the organizational infrastructure, legal authority, and governance structure to be able to comply with the obligations of the Common Agreement.
well as the DHCS Cal-HOP program’s concept of a Qualified Health Information Organization (QHIO).28,29

The state’s qualification processes and requirements should be overseen by CalHHS/the Center for Data Insights and Innovation (CDII) and developed through a transparent, public process. 


28 An HIO designated as “qualified” to participate in the Cal-HOP program and to receive payments for meeting milestones with qualified provider organizations. QHIO criteria included possession of specified organizational characteristics and technical capabilities as well as the ability to meet certain publication and reporting requirements. More information is available on the DHCS website.

Realizing the value of health information exchange requires not only the sharing of health and human service information but also its application and use to support individuals’ health and well-being. Event notifications – real-time, automated messages that can alert an individual's primary care practitioner, care manager, or care team of changes to the individual’s condition (e.g., admission to, discharge from, or transfer to [ADT] a health care facility) – are a primary example of how health and human service information exchange can be used to support care coordination and service delivery, though their utilization is often limited to a narrow set of participants and circumstances (e.g., alerting of transitions from acute care facilities).

Event notifications have the potential to improve care coordination and transitions among health and human service providers, thereby reducing the frequency of serious medical errors, delayed or inappropriate treatment, and prolonged lengths of stay.\(^{30,31}\) One 2017 study found that use of event notifications in the hospital setting resulted in a 2.9-percentage-point reduction in the likelihood of readmission.\(^{32}\)

Federal and state payers are increasingly recognizing the importance of providers connecting to data exchange intermediaries to send and receive event notifications (e.g., ADT alerts), particularly to support high-needs populations, and are incorporating such obligations into participation requirements and downstream contracts. In 2020, CMS published the Interoperability and Patient Access final rule, which included conditions of participation that required most hospitals to, by May 2021, send electronic ADT event notifications to all applicable post-acute care service providers and suppliers, in addition to the individual’s primary care practitioner, primary care group, or other practitioner or group that is identified by the individual that may be responsible for the individual’s care.\(^{33,34}\)


The federal interoperability rule, however, remains narrow in its requirements, limiting the potential scope of exchange and impact on individuals. For example, the final rule requirements only pertain to hospitals, do not require that notifications be sent to a member’s health plan, and only require that notifications be transmitted when triggered by inpatient and emergency department utilization. To realize the objectives of AB133, California should build upon federal requirements to expand the scope of ADT notifications, broadening the types of entities required to send data; the types of organizations the notifications should be sent to; and message content to include other human service-related transitions (e.g., housing, correctional facilities) that require notification.

Opportunity #3.1: Expansions to Federal Event Notification Requirements

**Summary:** The state should develop policy and contracting requirements that extend the scope of federal event notification requirements to include additional health and human service organizations and notification types.

The state should embrace and expand federal event notification requirements in California to other health and human service organizations involved in the delivery of care and services, including health plans and other human service providers such as housing agencies and correctional facilities. Expanding the scope of event notifications over time to include a broader set of entities would allow for a more complete understanding of an individual’s health and human service transitions and associated needs, and would support a whole-person approach to care. Broader event notification participation would be particularly beneficial for organizations serving and supporting high-needs individuals and those receiving support from multiple delivery systems and health and human service sectors.

The state should develop new requirements overseen by DxF Governance through a public and transparent process, proposing new event notification requirements and timelines that consider implementation feasibility. The state should also release guidance on best practices for how organizations of varying types may incorporate event notification alerts into practice, particularly for organizations new to sending or receiving notifications, and for use of new notification types (e.g., transitions from incarceration or from institutional settings).
Gap #4: Intra- and Inter-Sector Data Exchange Capabilities

**Relevant AB133 Provisions**

Identify gaps, and propose solutions to gaps, in the life cycle of health information, including gaps in:

- [the] storage, maintenance, and management of health information.  
  \[§130290(c)(3)(B)(iii)\]
- Linking, sharing, exchanging, and providing access to health information.  
  \[§130290(c)(3)(B)(iv)\]

On or before January 31, 2023, the California Health and Human Services Agency shall work with the California State Association of Counties to encourage the inclusion of county health, public health, and social services, to the extent possible, as part of the California Health and Human Services Data Exchange Framework in order to assist both public and private entities to connect through uniform standards and policies. It is the intent of the Legislature that all state and local public health agencies will exchange electronic health information in real time with participating health care entities to protect and improve the health and well-being of Californians.  
\[§130290(e)\]

California’s public health and human service agencies are stewards of valuable information that they need to manage programs and services they deliver to clients, individuals, and families. Most have mixed capabilities to electronically exchange timely and usable information with other health and human service organizations that could benefit from accessing it to provide a more complete picture and help them deliver a richer complement of coordinated services.

In California, county governments, in particular, have far-reaching responsibilities, partnering with the state to provide critical health care services to their residents. Responsibilities of California’s 58 counties include serving as the primary operator of health programs for low-income individuals without other forms of coverage and providing an array of behavioral health services, administered in part through county-operated mental health plans.\(^{35}\)

California’s state and local public health agencies also play a critical role in promoting and protecting public health, collaborating with health care organizations to carry out core public health activities, including disease surveillance and case reporting. However, despite long-standing working relationships, significant barriers to effective data exchange between public health and health care organizations remain. In 2019, 45% of Californian hospitals identified the inability of public health systems to receive

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data electronically as a major challenge to reporting health information. Researchers have cited underinvestment in public health infrastructure as a factor that has prevented more widespread public health data exchange across sectors, exacerbated by factors such as limited interoperability with health care EHRs and the challenges of onboarding and managing interfaces with multiple health systems.

With increasing recognition of the impacts of social factors on health, there is also growing support for the exchange of information about an individual’s social needs and the state and local government programs the individual may be participating in. However, efforts to increase data exchange between government entities, human service providers, and others involved in maintaining or improving health are complicated by operational, legal, and regulatory challenges for exchanging data across sectors (see Gap #E1 and associated opportunities).

Opportunity #4.1: Upgrades to California County HIT Infrastructure

**Summary:** The state should leverage and expand federally funded programs to upgrade state and local public health information technology infrastructure and to provide a glidepath for county health, public health, and human service entities to participate in information exchange.

The state should make strategic investments in state and county health, public health, and human service capacity to strengthen their participation in information exchange, as envisioned by AB133, the DxF, and the DxF DSA. California should prioritize the building of infrastructure to support data exchange and should develop an approach for ensuring the long-term sustainability of county technology infrastructure investments, identifying funding and resources to defray ongoing costs. Development and rollout of the capacity-building program would require close collaboration between state and local entities to leverage and align on priority objectives and initiatives.

The state should seek funding from federal sources to support upgrades to technology that can foster data sharing between state and county entities and stakeholder groups referenced in AB133. The program should leverage and align with federal modernization efforts, such as the Centers for Disease Control and Prevention (CDC) Data Modernization Initiative, which uses Coronavirus Aid, Relief, and Economic

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40 AB133 requires that specified entity types execute the DxF DSA. Specified entities include general acute care hospitals, physician organizations and medical groups, skilled nursing facilities, health service plans and disability insurers, Medi-Cal managed care plans, clinical laboratories, and acute psychiatric hospitals.
Security (CARES) Act funds to convene public health jurisdictions and other private- and public-sector entities to strengthen the nation’s public health data infrastructure.\(^{41}\)

**Opportunity #4.2: Public Agency Data Exchange Policy and Contracting Requirements**

**Summary:** The state should, through policy, procurement processes, and contract amendments, contractually obligate vendors to share information with health and human service organizations to advance goals envisioned by AB133.

The state should enact and support policies that expand health and human service data reporting and exchange requirements. Legislation pertaining to health and human service programs and initiatives could include provisions requiring applicable entities to share data to advance goals envisioned by AB133. For example, California’s AB977 requires that, beginning in January 2023, entities operating state homelessness programs report a set of specified data elements into their local Homeless Management Information System (HMIS) as a condition of receiving state funds.\(^{42}\) Data housed in regional HMIS systems is collected and warehoused in a statewide data system with the goal of supporting improved matching of data on homelessness to “programs impacting homeless recipients of state programs.”\(^{43}\)

In addition to enacting supportive legislation, the state could also include requirements for publicly funded programs to incorporate data sharing requirements into procurements and vendor contracts. Such policies would apply to use cases defined pursuant to the DxF and would include flow-down requirements for vendor contracting such as HMIS vendors and EHR vendors in correctional facilities. In one county-based example, Merced and San Joaquin counties have established contracts that require EHR vendors operating in their county jails to share health information with local data exchange intermediaries upon release of incarcerated individuals, to encourage coordinated transitions between the correctional facility and community settings.\(^{44}\)

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\(^{42}\) California Welfare and Institutions Code §8256(d).

\(^{43}\) California Welfare and Institutions Code §8257(b)(13).

B. Data Exchange Standards

Gap #1: Demographic and Human Service Data Exchange Standards and Capacity

**Relevant AB133 Provisions**

Identify ways to incorporate data related to underserved or underrepresented populations, including, but not limited to, data regarding sexual orientation and gender identity and racial and ethnic minorities. [§130290(c)(3)(D)]

Identify ways to incorporate data related to social determinants of health, such as housing and food insecurity, into shared health information. [§130290(c)(3)(C)]

Identify gaps, and propose solutions to gaps, in the life cycle of health information, including gaps in:

- Health information creation, including the use of national standards in clinical documentation, health plan records, and social services data. [§130290(c)(3)(B)(i)]
- Translation, mapping, controlled vocabularies, coding, and data classification. [§130290(c)(3)(B)(ii)]

Understanding who our systems of health serve and the barriers to health they confront is essential for ensuring the effective and equitable delivery of services to improve their lives and well-being. Demographic data – indicators of an individual’s race, ethnicity, sex assigned at birth, gender identity, sexual orientation, and spoken language, among other characteristics – provide health and human service organizations with critical information about the health and service needs of those they serve, and allow for the early identification and mitigation of access and service delivery inequities and health disparities. Social determinants of health (SDOH) data – including indicators of an individual’s human needs that expand upon the health care system (e.g., housing stability, food security, personal safety) and the human services they may utilize – similarly provide health and human service organizations insight into the health-determining needs of their populations, and where opportunities may exist to connect individuals to services to support their overall, whole person health and well-being.

However, the standardized collection, curation, and use of demographic and SDOH data in California remain uneven and developing. Strengthening demographic and SDOH data collection, curation, and use in our systems of health is critical for supporting California’s underserved and under-represented populations and addressing the health disparities they may confront.
Demographic Data

The COVID-19 pandemic illuminated the stark inequities in health service access and the devastating disparities in health outcomes that exist and persist among Californians, and reinforced the importance of collecting better and more usable demographic data about the populations and communities we serve. Demographic information available to health and human service providers is frequently incomplete and inaccurate, or is collected against varying standards, making its use challenging.

Race, ethnicity, and language data, for example, has traditionally been collected by health and human service programs and organizations on a voluntary basis, with little context provided for how such information would be used to support the individual, often resulting in low response rates. In California, a quarter of COVID-19 cases still have missing race and ethnicity data, two years into the pandemic.\(^45\) Analysis of national-level data similarly illustrates challenges that organizations face in collecting demographic information, with two-thirds of commercial health plans, half of Medicaid plans, and over one-quarter of Medicare plans reporting missing race data for at least half of their membership.\(^46\)

Different health and human service organizations collect race, ethnicity, and language information using different standards, making integration and comparability challenging. Even demographic data points with long histories of collection, such as gender, generally don’t reflect our current understanding of these attributes and aren’t collected with other contextual information (e.g., gender identity) to ensure that they may be meaningfully applied to support service delivery and health improvement. Federal standards for attributes like gender identity and sexual orientation are not widely implemented, and where information on these characteristics has been collected, it is frequently collected against local standards, which can vary significantly.

Social Determinants of Health (SDOH) Data

National and federally recognized standards for human service data exchange have not been broadly adopted, and the standardized collection, exchange, and use of information on SDOH remain limited. While health and human service organizations are increasing efforts to collect and use data on SDOH – which are estimated to

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account for approximately 80% of health outcomes\textsuperscript{47,48} – such efforts remain largely in their early stages.

SDOH data can be collected directly from interactions with an individual (e.g., patient screening, observed conditions noted in clinical records or on claims), but such data may not be collected in alignment with modern federal standards such as United States Core Data for Interoperability (USCDI) version 2, as some standards are relatively nascent.\textsuperscript{49} Use of SDOH-relevant ICD-10-CM codes in claims or encounters also remains limited, as health care organizations frequently do not have financial incentives or training to accelerate use.\textsuperscript{50,51}

SDOH data may also be collected indirectly from changes in an individual’s engagement with human service programs (e.g., CalFRESH enrollment, identification in HMIS). However, this data may not always be accessible in a timely manner to health care organizations that could use it to inform an individual’s care due to various legal, operational, and technical barriers to data exchange.\textsuperscript{52}

In addition to obstacles to data collection, health and human service organizations also face challenges with SDOH data exchange. At present, data exchange intermediaries, including national networks, regional HIOs, and EHR vendor networks, generally have limited capabilities to exchange structured and standardized SDOH and other human service data.\textsuperscript{53,54,55}

Standardized collection, exchange, and use of demographic and SDOH data can strengthen the ability of health and human service organizations to address the needs of individuals, improve the quality and cultural appropriateness of care delivery, and identify and reduce health disparities.


\textsuperscript{49} USCDI version 2 was released by ONC in July 2021. More information about USCDI is available \textsuperscript{HERE}.

\textsuperscript{50} SDOH-relevant IC-10-CM codes are primarily found within categories Z55–Z65 under Factors influencing health status and contact with health services. See CMS’ Official Guidelines for Coding and Reporting for more detail, available \textsuperscript{HERE}.


Opportunity #1.1: **Demographic Data Collection and Use Standards, Requirements, and Incentives**

**Summary:** The state should establish demographic data collection and sharing requirements and incentives through public and private payers to encourage data use among health and human service organizations in California.

The state should establish standards, requirements, and incentives to encourage the collection, exchange, and use of demographic data in California.

California may work with health and human service organizations to establish and require demographic data collection and reporting on data elements including, but not limited to, race, ethnicity, gender, sexual orientation, gender identity, and disability status. State standards should align with federal standards, such as USCDI version 2 and others such as those maintained by the U.S. Office of Management and Budget\(^{56}\) and the U.S. Centers for Disease Control and Prevention.\(^{57}\) California should also establish a process to evaluate relevant new federal standards as they are released, to determine need for implementation.

California should promote common standards across public and private organizations, learning from other state experiences. For example, in 2021, Connecticut passed legislation\(^ {58}\) that required:

- State agencies, boards, or commissions to collect demographic information to follow a set of collection and reporting standards;
- The collection of self-reported demographic information in EHRs; and
- That a state office be charged with evaluating standard race and entity categories in consultation with stakeholders.

The introduction of data standards may be paired with and reinforced by data collection, reporting, and use requirements by public and private payers in California. For example:

- DHCS is promoting increased demographic data collection to reduce disparities and inequities through CalAIM,\(^ {59}\) and is increasingly using available data to identify potential differences in health outcomes between racial and ethnic population groups, sharing findings with health plans to guide interventions.\(^ {60}\)
- Covered California continues to increase its race and ethnicity data reporting and use expectations, requiring Qualified Health Plans (QHPs) to achieve 80% self-
identification of race and ethnicity data for their Covered California enrollees and to work with Covered California to reduce identified disparities.\textsuperscript{61}

Nationally, CMS has prioritized its expansion of the “collection, reporting and analysis of standardized data” – including race, ethnicity, language, sexual orientation, gender identity, and disability status – in its Medicare Equity Plan,\textsuperscript{62} as broader federal action in this area has been encouraged.\textsuperscript{63}

Health and human service organizations subject to new data collection, reporting, and use requirements may require technical assistance to support implementation.

Opportunity #1.2: Social Determinants of Health (SDOH) Data Collection and Use Standards, Requirements, and Incentives

**Summary:** The state should establish SDOH data collection and sharing requirements and incentives through public and private payers to encourage data use among health and human service organizations in California.

The state should adopt federal SDOH data standards and should establish requirements and incentives to encourage data collection, exchange, and use in California, helping health and human service entities better understand individuals’ broad health and human service needs – and the barriers they might confront in addressing them.

Specifically, the state should evaluate and adopt USCDI version 2 standards, which include standards pertaining to SDOH care goals, assessments, health concerns, and interventions.\textsuperscript{64} Widely used among health care organizations, USCDI is a set of health data classes and elements developed by the U.S. Office of the National Coordinator for Health Information Technology that supports standard, nationwide, and interoperable health information exchange. California should also establish a process to evaluate new federal standards as they are released (e.g., USCDI version 3) for potential implementation over time.\textsuperscript{65}

The adoption of statewide SDOH data standards may be paired with common data collection, data quality, and performance reporting requirements and incentives through public and private payers to support system change. To inform state approaches,


\textsuperscript{65} ONC is currently developing USCDI version 3, with an expected release date in July 2022. The draft version of USCDI version 3, which was released for public comment, retains the SDOH-related elements contained in version 2. More information is available on the ONC website HERE.
California should examine the experiences of other states that have incorporated SDOH screening and referral requirements in their Medicaid managed care contracts and used quality reporting requirements and incentive payments to monitor and reward progress in meeting SDOH-related goals.66 Health and human service organizations subject to requirements or incentives pertaining to SDOH data collection, exchange, and use practices may require technical assistance to support implementation.

Establishing statewide SDOH data collection, exchange, and use requirements and incentives would build upon significant system investments in California and would position California as a leader among states advancing whole person health and health equity. In California, the Department of Health Care Services is reshaping its Medi-Cal program and delivery system through an expansive reform effort – CalAIM – prioritizing the identification and mitigation of social needs for high-risk, high-utilizing enrollees through increased service integration among counties, health plans, and community-based organizations (CBOs).67 Nationally, CMS has encouraged state Medicaid and Children’s Health Insurance Program (CHIP) initiatives to adopt strategies like CalAIM that address SDOH, including instituting new data collection and reporting requirements, investing in systems capable of exchanging SDOH data, and establishing programs that can put such data to meaningful use.68

Opportunity #1.3: Cross-State Agency Data Sharing Enhancements

**Summary:** The state should strengthen data exchange relationships between state agencies, supporting the exchange of program enrollment and relevant demographic and SDOH information to support a coordinated approach to health care and human service delivery across disparate programs and settings.

The state should enhance cross-agency program enrollment, service utilization, demographic, and SDOH data sharing, in alignment with state and federal law, to support cross-program service coordination and greater shared understanding of population health and human service needs. The state should also permit access to this data by external health and human service organizations that are serving Californians, to the extent allowable by state and federal law, in order to improve service delivery.

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66 States pursue a number of strategies to advance the collection, exchange, and use of SDOH information, including requirements for entities to screen individuals for social needs or to capture SDOH data using ICD-10-CM “Z” codes. More information can be found in a Kaiser Family Foundation brief describing Medicaid Authorities and Options to Address SDOH, available [HERE](https://www.kff.org/medicaid/issue-brief/medicaid-authorities-and-options-to-address-social-determinants-of-health/). 

67 Department of Health Care Services, CalAIM, accessed April 28, 2022, [https://www.dhcs.ca.gov/CalAIM/Pages/calaim.aspx](https://www.dhcs.ca.gov/CalAIM/Pages/calaim.aspx).

Examples of the types of state agency data that may be shared to support coordinated service delivery are shown in Table 1.69

Table 1

<table>
<thead>
<tr>
<th>State Government Entity</th>
<th>Relevant Programs or Topics (Example)</th>
<th>Relevant Data Systems (Example)</th>
<th>Data Types (Example)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Department of Health Care Services (DHCS)</td>
<td>Medi-Cal</td>
<td>Medi-Cal Eligibility Data System (MEDS)</td>
<td>Program enrollment and identified social needs (if captured on claims or encounters)</td>
</tr>
<tr>
<td>California Department of Social Services (CDSS)</td>
<td>CalFresh, Child Welfare</td>
<td>California Statewide Automated Welfare System (CalSAWS)</td>
<td>Program enrollment and other food insecurity indicators</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Child Welfare Services/Case Management System (CWS/CMS)</td>
<td>Child welfare involvement indicators and identified health and social needs</td>
</tr>
<tr>
<td>Business, Consumer Services and Housing Agency (BCSH) – California Interagency Council on Homelessness (Cal ICH)</td>
<td>Homelessness</td>
<td>Homeless Data Integration System (HDIS)70</td>
<td>Housing services enrollment and other housing stability indicators</td>
</tr>
<tr>
<td>Attorney General’s Office</td>
<td>Criminal Justice</td>
<td>Criminal Justice Statistics Center</td>
<td>Justice involvement indicators including information on probation events and transitions</td>
</tr>
<tr>
<td>California Department of Developmental Services (CDDS)</td>
<td>Developmental Disabilities</td>
<td>SANDIS</td>
<td>Diagnostic and evaluative information</td>
</tr>
<tr>
<td>California Department of Public Health (CDPH)</td>
<td>Public Health</td>
<td>California Immunization Registry (CAIR)</td>
<td>Public health programmatic data (e.g., immunizations, case and lab reports)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>California Reportable Disease Information Exchange (CalREDIE)</td>
<td></td>
</tr>
</tbody>
</table>

69 Table 1 provides examples of entities and data types that may be involved in efforts to enhance cross-agency data sharing. The list of entities and data types is not intended to be understood as comprehensive.

70 The Homeless Data Integration System (HDIS) is a statewide data warehouse that compiles and processes data from all 44 local homelessness response systems in California. More information on HDIS is available on the BCSH website [HERE](#).
As California continues to scale programs that support whole person care, the state may evaluate incentivizing health and human service organization connections to health information exchanges, community information exchanges (CIEs), and other data exchange intermediaries to support cross-program and cross-sector exchange of program, service utilization, and individual demographic and SDOH data.

Investments in cross-state agency data sharing may require the state to identify new resources – including sources of funding – to support implementation. Each agency should investigate, identify, and evaluate federal funding opportunities to support initiative goals. For example, DHCS should explore the possibility of leveraging enhanced federal Medicaid matching funds, as described in CMS guidance, for “state expenditures to design, develop, install, or enhance” interoperable systems capable of identifying individuals’ SDOH needs and sharing that information with appropriate medical and social support services.71 (See Opportunity #F1.1 for more information.)

Opportunity #1.4: Recommendations to the Federal Government to Improve Demographic and SDOH Data Collection Nationally

**Summary:** The state should develop and advance recommendations to the federal government to improve demographic and SDOH data exchange standards nationally.

The state should advance recommendations to the federal government to improve demographic and SDOH data collection nationally. The collection of demographic and SDOH data remains uneven across health and human service organizations, with variation in data standards – and data standards implementation – even among federal agencies.

While it seeks to align with federal requirements, California should develop an approach to advocate for and advance standards that reflect local learnings and priorities. The approach should include a process to identify opportunities to comment on national data standards, including those promulgated by federal agencies such as ONC, CMS, the Health Resources and Services Administration (HRSA), the U.S. Department of Health and Human Services Office for Civil Rights (OCR), and the Substance Abuse and Mental Health Services Administration (SAMHSA), as well as those proposed by non-governmental organizations such as the American Medical Association; the University of California - San Francisco/Social Interventions Research and Evaluation Network-supported The Gravity Project; and the Sequoia Project. For instance, California – through its Center for Data Insights and Innovation – could collect and advance recommendations to the federal government to address misalignment between HRSA’s demographic standards and those of the USCDI, which creates reporting burden for

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health care providers that may have to report to different payers using differing standards.

The state should also develop processes to assess whether new proposed federal standards and regulations should be incorporated into the DxF and its DSA, and should potentially provide guidance to implicated health and human service organizations on how they may effectively implement these standards and regulations. For example, CalHHS could develop a consolidated response to the emerging draft USCDI version 3 standards, which support more consistent functional, cognitive, and other disability status data collection, sharing, and use, to reflect local learnings and potentially improve proposed standards before they are finalized.
C. Provider Information and Person Identity Management

Gap #1: Robust Provider Information

Relevant AB133 Provisions
Identify gaps, and propose solutions to gaps, in the life cycle of health information, including gaps in linking, sharing, exchanging, and providing access to health information. [§130290(c)(3)(B)(iv)]

Collection and management of provider information is critical for identifying the methods, standards, and routes for exchanging health and human service information to support care coordination and other DxF use cases. However, complete provider information (e.g., name, practice locations, organizational affiliates, and the available means to exchange health information such as Direct addresses and endpoints) is not always available or accessible to all health and human service organizations across California, creating barriers to effective information exchange and service delivery.

Provider identities are often assigned and/or managed by federal, national, or state provider directories, the scopes and utility of which may vary based on their purpose. Provider directories may be established for more consumer-facing purposes (e.g., helping individuals identify in-network providers close to where they live or work) or to support provider-to-provider communications (e.g., listing the means, data exchange intermediaries, and addresses available to send data to or request data from a provider). Examples of provider directories used to support provider-to-provider communications in California include those maintained by the California Trusted Exchange Network (CTEN), Carequality, DirectTrust, and the National Plan & Provider Enumeration System (NPPES), which is the CMS program responsible for assigning National Provider Identifiers (NPIs).

Federal and state policies require and govern the use of provider directories for a limited set of use cases. For example, CMS’ recent Interoperability and Patient Access final rule and California’s SB137 require that certain payers maintain or make available provider directory information. The CMS Interoperability and Patient Access final

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rule also separately encourages providers to maintain accurate digital contact information that can be used to exchange health information electronically.\textsuperscript{77}

Despite their importance, provider directories are often limited in their ability to support many of the use cases envisioned by AB133 and the DxF and its DSA. For example:

- Participation in provider directories is often voluntary or limited to the participants of a given network, limiting the directories’ scope and ability to support communication and information exchange between disparate entities and sectors.\textsuperscript{78}
- Many provider directories are not designed for the purpose of facilitating health information exchange and lack critical digital contact information that would facilitate electronic exchange.
- Provider directories are only as useful as the timeliness of their data, and as provider contact and system affiliation information can often change, protocols and incentives or regulatory requirements must be in place to ensure that providers are reflecting those changes in the directory.

Basic provider information and accurate information about how to exchange health and human service information with providers are critical for creating a shared understanding of the health and human service organizations supporting Californians’ health and well-being, knowing whom those organizations are serving and for what services, and connecting health and human service organizations as they strive to deliver more coordinated and whole person care. California presently lacks a comprehensive provider directory strategy capable of advancing these objectives.

**Opportunity #1.1: Provider Directory and Reporting**

**Summary:** The state should support the establishment of a statewide provider directory and should require signatories of the DxF DSA to contribute data to encourage provider-to-provider communication and information exchange.

The state should support the establishment of a statewide provider directory and should develop supporting policies to encourage communication and information exchange between disparate health and human service entities. The provider directory may be newly established, or may leverage and build upon existing directories and approaches, and should be designed for the purposes of enabling provider-to-provider communication. The provider directory should include, among other identifying data elements, information on provider digital contact information for various existing industry


\textsuperscript{78} As an example, CMS encourages providers to report digital contact information in NPPES but does not levy financial penalties or restrictions on providers that fail to do so. More information on digital contact information in NPPES is available HERE.
standards (e.g., Direct messaging address, Fast Healthcare Interoperability Resources® [FHIR®] URL).

All signatories of the DxF DSA should be responsible for submitting required data elements, including digital contact information, to the provider directory using a standardized process, as well as for supporting the maintenance and timely updating of such information on an ongoing basis. DxF Governance should consider technical approaches and specifications to support provider identification and data exchange and should clarify requirements and guidance for DxF participants in the DxF DSA and P&Ps.
Gap #2: Person Identity Management

**Relevant AB133 Provisions**

On or before July 31, 2022, the California Health and Human Services Agency shall develop in consultation with the stakeholder advisory group in subdivision (c) a strategy for unique, secure digital identities capable of supporting master patient indices to be implemented by both private and public organizations in California. [§130290(h)]

The effective exchange and use of health care and human service data to support treatment and care is dependent upon linking data to the right individual. Many health care providers, health plans, and data exchange intermediaries have robust person resolution and record-linking technologies within their organizations. However, no robust or systematic coordination of digital identities, person resolution, or data linking exists across organizational boundaries in California, limiting the efficacy of cross-organizational data exchange.

As a result, organizations often fail to locate existing health records for individuals they serve that might exist at other organizations to support care coordination and management, because the organizations’ health information systems fail to agree on a single personal “identity.” Conversely, records for different individuals may be inappropriately matched, presenting an inaccurate picture of a person’s health and human service information – and potentially risks to health, if incorrectly linked data is used to guide clinical care. Both issues can lead to delays in effective service provision, poorer health outcomes, or even errors in service delivery.

**Opportunity #2.1: Strategy for Digital Identities**

**Summary:** The state should adopt the Strategy for Digital Identities called for in AB133 as a component of the DxF.

California stakeholders have significant experience in person resolution, person matching, and record linking through participation in existing networks. This experience was leveraged to create a Strategy for Digital Identities that focused on linking health and human service information to a real person across organizational and sector boundaries.

*For more details, please defer to the Strategy for Digital Identities document, available at the following link.* [final link to be inserted]
D. Individual Data Access

Gap #1: Individual Data Access

**Relevant AB133 Provisions**

Identify gaps, and propose solutions to gaps, in the life cycle of health information, including gaps in linking, sharing, exchanging, and providing access to health information. [§130290(c)(3)(B)(iv)]

Identify how all payers will be required to provide enrollees with electronic access to their health information, consistent with rules applicable to federal payer programs. [§130290(c)(3)(I)]

Individuals consistently face challenges in accessing and contributing to their health and human service records – directly or through selected third parties – in a manner that is convenient, timely, and compliant with federal access requirements.

Under the federal Health Insurance Portability and Accountability Act (HIPAA), individuals have a legal, enforceable right to access their health records maintained by a covered entity such as a provider or health plan. Research has shown that providing individuals access to their own health information is associated with improvements in doctor-patient communication, treatment adherence, and individual empowerment – and is of great interest to many individuals. According to a national survey conducted in 2020, the majority of respondents reported they would want to access their own health information, with particularly high interest in accessing laboratory test results (89%) and history of medical conditions (88%).

Despite individuals’ right to access their information and their interest in doing so, numerous barriers to access exist, including complex access requirements; fragmented data; financial costs (e.g., printing fees); and, often, health care organization non-compliance. Even for individuals who are able to access their information, their ability to contribute data – that is, to engage in bidirectional exchange – is often limited despite its potential benefits, such as facilitating submission of patient-reported outcomes or even corrections to errors in an individual’s medical record.


Recent policy has taken steps to support data access for individuals, including through ONC’s 21st Century Cures Act final rule, which includes provisions that aim to curtail information blocking, and CMS’ Interoperability and Patient Access final rule, which requires CMS-regulated payers to make individuals’ data, including claims and encounter data, available to individuals via a secure, standards-based Application Programming Interface (API). While these federal rules have the potential to advance information access for individuals, barriers remain, including stakeholder concerns regarding data confidentiality and privacy, which have surfaced and been heightened as a result of the push for more open exchange of data.

Opportunity #1.1: Policies to Ensure That Individuals Have Meaningful Access to Their Longitudinal Health Information

**Summary:** The state should adopt and expand policies to ensure that individuals have meaningful access to their longitudinal health information across all health care organizations that are required to execute the DxF DSA, allowing all Californians to be meaningfully engaged in their care and make informed health care decisions.

The state should adopt policies to ensure that individuals have meaningful access to their longitudinal health information across all health care organizations subject to AB133, allowing them to make informed health care decisions.

Potential policies and requirements to advance this objective may include building upon federal rules that increase individuals’ access to their health information to include all health care organizations subject to the DxF and its DSA. Specifically, the state could expand on ONC’s Cures Act final rule to apply the federal information-blocking provisions to the state’s licensed health plans and health care organizations required to execute the DxF DSA. The state could also expand on CMS’ Interoperability and Patient Access final rule to require that signatories of the DxF DSA implement and maintain a secure, standards-based API (e.g., Health Level 7® (HL7®) FHIR® Release 4.0.1) that allows individuals to easily access their health information through third-party applications of their choice.

The state should also evaluate where California law governs individuals’ access to their longitudinal health information and may need to be modified, as federal rules often defer to state law. Examples of relevant state law include California Health and Safety Code §130290(f)(1-6).
123110, which gives individuals the right to inspect and receive a copy of their patient records by request and upon payment of any reasonable costs.87

Supportive policies may also provide individuals with bidirectional access to their health information maintained by health care organizations – including to add self-reported health information and to correct inaccurate information to the extent allowed by law – in a manner that accommodates their preferred language, readability level, and disability status while also supporting access for those Californians who have limited access to technology or broadband or have low health and information technology literacy.

Health care organizations should be expected to provide individuals with education and support on how to access, update, and use their health information to inform their decision-making.

Opportunity #1.2: Policies to Strengthen Understanding and Trust for Individuals Sharing Data

**Summary:** The state should assess, strengthen, and adopt policies that will support individuals in understanding how their health and human service information may be used and will help establish trust in the systems in place that govern its use.

The state should adopt policies to ensure that individuals understand how their health and human service information may be used and to build trust in the systems in place that govern its use.

California should review its fair information practices and policies, consider updates, and develop stronger guidelines for data use that can be linked with the DxF DSA and referenced in future state policy guidance. Guidelines may include and address:

- Transparent data policies;
- Limitations on health information collection, use, and disclosure;
- Consent procedures;
- Data quality, integrity, and security protections; and
- Accountability processes through appropriate audit trails and enforcement.

Guidelines should be clear on how individuals’ health and human service information may be shared and where individuals have the ability to provide or withhold consent prior to its exchange and use, in alignment with state and federal laws. The U.S. Department of Health and Human Services Office of Civil Rights’ guidance on the right

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87 California HSC §123110, available [HERE](#).
of individuals under HIPAA to access their health information\textsuperscript{88} and ONC’s 2018 Model Privacy Notice\textsuperscript{89} provide examples of clear guidance to inform individuals about data access and privacy policies.

California should develop practices and policies to hold DxF DSA participants accountable for participation in the appropriate exchange and use of individuals’ health information, including by monitoring and evaluating participation and adherence to policies regarding the sharing, safeguarding, and use of health information; integrating explicit protections against misuse of health data; supporting health care organization training on individuals’ access to health information; and supporting mechanisms to solicit and process feedback from individuals to improve data access.


E. Data Exchange Law, Regulations, and Policy

Gap #1: Data Exchange Law, Regulations, and Policy

**Relevant AB133 Provisions**

The California Health and Human Services Data Exchange Framework shall align with state and federal data requirements, including...applicable state and federal privacy laws related to the sharing of data among and between providers, payers, and the government, while also streamlining and reducing reporting burden. [$130290(a)(3)]

Address the privacy, security, and equity risks of expanding care coordination, health information exchange, access, and telehealth in a dynamic technological, and entrepreneurial environment, where data and network security are under constant threat of attack. [$130290(c)(3)(F)]

Identify ways to incorporate relevant data on behavioral health and substance use disorder conditions. [$130290(c)(3)(E)]

Identify gaps, and propose solutions to gaps, in the life cycle of health information, including gaps in linking, sharing, exchanging, and providing access to health information. [$130290(c)(3)(B)(iv)]

Numerous federal and state laws, regulations, and policies that govern the exchange of physical, behavioral health, and human service data create real or perceived barriers to sharing information that is necessary to inform whole person care and population health goals.

Legal protections safeguard the privacy and security of personal information. The federal Health Insurance Portability and Accountability Act is a foundational regulation that governs the disclosure of protected health information (PHI), setting national standards for its collection, exchange, and use.

However, many data types created and used by health and human service organizations are also protected by myriad other federal and state laws, regulations, and policies. For example, certain data types, including those pertaining to behavioral health, HIV/AIDS, sexual health, and minors, are governed by specific federal and state rules and regulations that require individuals’ authorization to disclose information for data sharing purposes. Beyond the domain of clinical health care, the collection and exchange of data curated by human service organizations or government agencies – including data related to housing, food security, and criminal history – may require
separate, individual authorization to be shared and are often protected by additional regulations and rules specific to the data collecting entity and data type.\textsuperscript{90,91}

While critical to ensuring appropriate stewardship of personal information and building trust among exchange partners, legal protections for subsets of health and human service data can present barriers to meaningful data exchange. In a study of California’s Whole Person Care (WPC) pilots, 16 pilots (64\%) reported patient privacy and confidentiality regulations as a major implementation challenge.\textsuperscript{92} Some pilots additionally noted that even when legal barriers were eliminated, fear and risk aversion affected organizations’ willingness to exchange data.

Effective data sharing requires health and human service organizations to have a common understanding of when information sharing is permissible and to have the capability to obtain, manage, and share information about an individual’s authorization to release and share data. However, managing authorization and consent can be challenging, with organizations’ operational approaches – and the effectiveness of these approaches – varying widely, hampering efforts to share authorizations across organizational boundaries.

**Opportunity #1.1: “Universal” Release-of-Information Authorization Form**

**Summary:** The state should support the adoption and use of a “universal” release-of-information authorization form to enable standardized data exchange in support of state priority use cases and the goals of AB133.

The state should support the adoption and use of a “universal” release-of-information authorization form. Universal authorization forms are characterized by their standardized structure and broad acceptance and use by disparate organizations, addressing legal-interpretation inconsistency and operational/review inefficiency to enhance trust and transparency between exchange partners and individuals. Broad-based adoption and use of universal authorization forms may improve the ability of individual organizations to collect and manage authorizations, reduce administrative burden, and lower barriers to data sharing across entities and sectors.

The scope of the universal authorization form should be determined in alignment with state priorities and the requirements of the DxF and its DSA. The state should develop the form to prioritize data types where authorization collection and sharing challenges are most acute and to prioritize use cases related to state investments in health and


human services, including applications stemming from CalAIM, the public health emergency, and other priorities. The state should develop the form leveraging lessons from other states and California’s WPC pilots. For example, in 2015, the Michigan Department of Health and Human Services (DHHS) released a standard consent form for sharing behavioral health information, the Behavioral Health Standard Consent Form, to address challenges with sharing such data. In 2018, Michigan DHHS released a policy bulletin specifying that the form must be “accepted, honored, and used” by all Medicaid providers that are requesting release of behavioral health-related information, in cases when such consent is required. California may also use policy levers to require or encourage use of the universal authorization form to maximize its benefits. In Michigan, for example, the state Legislature required that all entities involved in treating those with certain behavioral health conditions honor and accept the form unless held to more stringent protections under federal law.

California should provide technical assistance to support organizations in understanding how and when to use the final authorization form to build knowledge as well as trust between users. Technical assistance should leverage existing state resources, such as the State Health Information Guidance (SHIG), and should provide organizations with support for collecting, using, and exchanging sensitive information, including that which may be specially protected.

Opportunity #1.2: Consent Management Service

Summary: The state should consider support for a consent management service that would allow individuals to electronically manage – and health and human service organizations to access – consents to disclose and share certain types of health and human service information.

The state should consider support for establishing a consent management service that would allow individuals to electronically provide, revoke, and manage their consent for access and use of their health and human service information. Health and human service organizations would then be expected to access the consent management service to validate whether – and with whom – they are permitted to exchange applicable data types for individuals under their care. A statewide service could

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improve care coordination and continuity for individuals and reduce the burden on individual health care entities to obtain and manage authorizations to share data.

The design and structure of a potential consent management service should be aligned with state priorities and the requirements of the DxF and its DSA. The state should consult with stakeholders to develop an overarching approach for the oversight and management of a potential service to include identifying the data types, types of consent, and scope of health care organizations the service would support as well as determining the entities that would be expected to participate.

The state should expect to provide technical assistance to health and human service organizations to support the transition to electronic modes of consent management, including providing materials that organizations can use to educate individuals on the purpose of the system, building trust, communicating the benefits of the system, and ensuring that the design of a potential service prioritizes individual privacy rights.

Consideration of a consent management service should be pursued in concert with other aligned opportunities. Specifically, strong person identity management, as described in the DxF Strategy for Digital Identities (see Opportunity #C2.1), is a prerequisite for effective consent management that crosses organizational boundaries. Additionally, development of a universal authorization form (see Opportunity #E1.1) would support the standardization of consent information, laying the groundwork for the design and implementation of an electronic consent management service that would be available and useable by disparate entities across the health and human service sectors.
F. Health and Human Service Information Exchange Financing

Gap #1: Financing to Support Health and Human Service Information Exchange

**Relevant AB133 Provisions**

Identify federal, state, private, or philanthropic sources of funding that could support data access and exchange. [§130290(c)(3)(K)]

Many barriers to data exchange are rooted in challenges for health and human service organizations in accessing financing to make needed health information technology investments. These barriers include but are not limited to:

- The limited adoption of EHRs and other technology capable of sharing information electronically among health and human service organizations (Gap #A1);
- Challenges with onboarding to qualified data exchange intermediaries (Gap #A2); and
- Insufficient data infrastructure in state and local public health departments (Gap #A4).

Funding is needed to establish an environment in California that is supportive of health and human service data exchange and to provide direct support for the organizations participating in exchange, particularly those that are under-resourced. Research suggests that financing programs can, when carefully considered and well implemented, meaningfully improve data exchange.97

While some federal and state investment opportunities – including HITECH-funded EHR incentive programs98 and Cal-HOP99 – have recently concluded, many federal, state, private, and philanthropic funding opportunities remain that should be leveraged to overcome identified gaps.

**Opportunity #1.1: Leveraging Governmental, Private, and Philanthropic Sources of Funding**

**Summary:** The state should identify and pursue federal, state, private, and philanthropic funding opportunities to finance data exchange priorities consistent with the vision and goals of the DxF and AB133.

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To support robust exchange of health and human service information in California, the state should identify, pursue, and leverage federal, state, private, and philanthropic funding opportunities. These funding opportunities, which vary in intended purpose and amount of funding available, should be assessed and pursued to finance data exchange priorities consistent with the vision and goals of the DxF and AB133. Potential funding opportunities that California should assess and consider leveraging are described below.

**CMS Medicaid Enterprise Systems (MES) Enhanced Funding**

States are eligible to receive enhanced federal funding for activities related to their Medicaid Enterprise Systems (MES), an umbrella term that refers to states’ Medicaid eligibility and enrollment (E&E) systems as well as Medicaid Management Information Systems (MMIS), the state IT infrastructure that supports Medicaid program management and administrative functions such as claims processing, beneficiary and provider management, clinical decision support, care management, and program integrity. MES enhanced funding may be used to support Medicaid systems initiatives related to claims, enrollment, health information exchange, and public health such as the development of master person indexes, provider directories, and other functions that allow Medicaid providers, beneficiaries and other users individuals to access claims, clinical or other health information.

States can receive a 90% federal match for system design, development and installation activities including planning activities, building interfaces and establishing connectivity, and initial software leasing or licensing. States can receive a 75% federal match for activities related to maintenance and operations such as system or software maintenance and ongoing software leasing or licensing. The federal share of funds may also depend on the cost allocation methodology that is proposed by states and accepted by CMS. The methodology can vary based on the use case or module (e.g., based on Medicaid covered lives as a percentage of state population, based on Medicaid providers as a percentage of all providers).

To request federal funding for MES initiatives, state Medicaid agencies submit planning, implementation, and operations advance planning documents (APDs) to CMS describing the funding need and proposed use of federal dollars and demonstrating compliance with applicable conditions and standards. CMS has emphasized the importance of reusing existing information technology assets and taking an incremental,

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100 42 C.F.R. 433.111(b)(1).
modular approach in state projects seeking enhanced funding, with the goals of reducing costs, mitigating project risk, and improving collaboration between states.\textsuperscript{104,105}

**CMS Medicaid Matching Funds**

States can pursue alternative mechanisms to access federal Medicaid matching funds to support activities related to the adoption and use of information exchange technologies when such activities may not be eligible or otherwise appropriate for MES enhanced funding. States can leverage matching funds as determined by the Federal Medicaid Matching Rate (FMAP), which determines the federal share of the cost of providing needed services to Medicaid beneficiaries.\textsuperscript{106} California’s FMAP in federal fiscal year 2023 is expected to be 50%.\textsuperscript{107} Medicaid administrative costs are funded separately but typically also at a 50% federal match rate, with some exceptions. States have leveraged these non-MES matching funds to support information exchange priorities. Arizona, for example, has leveraged federally matched dollars to establish a data sharing incentive program that increases payments for eligible entities that share data with Health Current, the state’s designated health information exchange.\textsuperscript{108}

**CMS Medicaid Waivers**

Under sections 1115 and 1915 of the Social Security Act, states can seek CMS approval of waivers of certain federal Medicaid requirements to allow for state flexibility in the use of federal funds to support Medicaid data infrastructure and exchange. For example, California has recently received authorization under a Section 1115 waiver for Providing Access and Transforming Health (PATH), a five-year, $1.85 billion initiative to support infrastructure and capacity building, including the strengthening of data exchange capacity in the Medi-Cal delivery system.\textsuperscript{109}

**CDC Section 317 of the Public Health Service Act**

States access federal funding through Section 317 of the Public Health Service Act to advance public health data infrastructure priorities. The Section 317 Program, enacted in 1962, is administered by the CDC and provides flexible funding for immunization program operations and infrastructure as well as vaccine purchase at the local, state, and national levels. Program funding can be used toward the development and


\textsuperscript{107} FMAPs for all states will be increased by 6.2 percentage points if the public health emergency is renewed October 2023. In this scenario, California’s FMAP would be 56.2%. Source: Kaiser Family Foundation.


implementation of immunization information technologies. For example, funding has
been used to enhance interoperability of public health systems with the clinical systems
used by other health care organizations as well as to help develop reminder and recall
systems that send providers and individuals a notification when the individual is due or
past due for a vaccination.\footnote{National Vaccine Advisory Committee, “Protecting the Public’s Health: Critical Functions of the Section 317 Immunization Program — A Report of the National Vaccine Advisory Committee,” Public Health Reports 128, no. 2 (2013): 78–95. https://doi.org/10.1177/003335491312800203.} The Section 317 Program is a discretionary program, and thus funding varies from year to year subject to the annual appropriations process; the
program was provided an appropriation of $651 million for federal fiscal year 2022.\footnote{The 317 Coalition - FY 2023 Labor-HHS Appropriations Bill,” The 317 Coalition, accessed May 11, 2022, https://www.317coalition.org/_files/ugd/cbc5b5_800ce7cc5b84350b8e48e593041bf16.pdf.}

State Taxes

States can also establish taxes to fund data exchange priorities. Taxes, which may be
broad-based or targeted, can serve to establish a state-level funding source that can
help augment — and in some cases, draw down — federal financing that can be used to
advance data infrastructure and exchange. For example, California’s Mental Health
Services Act (MHSA), passed by voters in 2004, is funded by a 1% income tax on
personal income in excess of $1 million per year and is used to finance a broad set of
programs and priorities that support the public behavioral health system, including
infrastructure and technology.\footnote{Mental Health Services Act, California Department of Health Care Services, accessed May 11, 2022, https://www.dhcs.ca.gov/services/MH/Pages/MH_Prop63.aspx.}

Federal, Private, and Philanthropic Grants

States can also leverage other federal funding sources — in addition to those previously
mentioned — as well as grants from private and philanthropic sources to blend and braid
funding to finance investments in data infrastructure and capacity across the state.
Federal agencies, including the HRSA and the Federal Communications Commission,
support an array of objectives foundational to data exchange, such as enhanced
funding opportunities that provide funds directly to delivery system organizations (e.g.,
the U.S. Department of Housing and Urban Development’s annual discretionary funding
allocations to Continuum of Care programs that address homelessness,\footnote{Continuum of Care (COC) Program Eligibility Requirements,” HUD Exchange, U.S. Department of Housing and Urban Development, accessed April 29, 2022, https://www.hudexchange.info/programs/coc/coc-program-eligibility-requirements/.} the
Administration for Community Living’s funding in support of Aging and Disability
Networks\footnote{Program Areas,” Administration for Community Living, accessed April 29, 2022, https://acl.gov/programs.}}