Charting a Course for an Equity-Centered Data System:

Recommendations from the National Commission to Transform Public Health Data Systems

OCTOBER 2021
The National Commission to Transform Public Health Data Systems was convened by The Robert Wood Johnson Foundation.

EDITORS:

**GAIL C. CHRISTOPHER**, DN, Commission Director and Executive Director, National Collaborative for Health Equity

**EMILY B. ZIMMERMAN**, PhD, MS, MPH, Associate Professor, Virginia Commonwealth University, Department of Family Medicine and Population Health, Division of Epidemiology; Senior Researcher and Director of Community Engaged Research, Virginia Commonwealth University Center on Society and Health

**ANITA CHANDRA**, DrPH MPH, Vice President and Director of RAND Social and Economic Well-Being, RAND Corporation

**LAURIE T. MARTIN**, ScD MPH, Senior Policy Researcher, RAND Corporation

Robert Wood Johnson Foundation

The National Commission to Transform Public Health Data Systems was convened by The Robert Wood Johnson Foundation.
# Table of Contents

A Statement from Dr. Richard E. Besser 2

Summary of Recommendations 3

Call to Action Summary 4

Introduction 6

Members of the independent National Commission to Transform Public Health Data Systems 10

Recommendations 11

Center Health Equity and Well-Being in Narrative Change 12

Prioritize Equitable Governance and Community Engagement 16

Ensure Public Health Measurement Captures and Addresses Structural Racism and Other Inequities 25

Commission Formation and Deliberative Process 32

Why this Commission Now? 33

Preparation 33

Selection of Commission Members and Deliberation Process 36

Conclusion 39

Call to Action 40

Acknowledgements 44

Glossary 44

References 46

Resources 48

Meet the Commissioners 49
A Statement from Dr. Richard E. Besser

Public health data should reflect the needs of everyone in America—regardless of their race, where they live, or how much money they have. If we are to become a nation that truly values health equity, public health data must reflect fairness, justice, and inclusion as fundamental requirements.

That's why, as the nation's largest health philanthropy, the Robert Wood Johnson Foundation (RWJF) convened the first-of-its-kind National Commission to Transform Public Health Data Systems. The Commission was charged with reimagining how data are collected, shared, and used, and identifying what public and private sector investments are needed, with the express purpose of improving health equity.

The COVID-19 pandemic laid bare the deficits of our public health infrastructure. It forced us to acknowledge that we must modernize our data systems—and we must act fast. Due to incomplete and disparate data collection, it took far too long to see the disproportionate impacts of the pandemic on Black, Latino, and Indigenous communities. Lack of reliable data collection on factors that matter for health equity and lack of consistent and transparent reporting of these data impeded us from saving lives and affected our ability to ensure that those hardest hit by COVID-19 were prioritized at every stage of the pandemic—from testing to hospitalization to vaccination.

The U.S. Centers for Disease Control and Prevention, along with more than 200 state and local jurisdictions, have declared racism a public health crisis. Reforming our public health systems to address structural racism requires that we collect and analyze data in ways that accurately reflect the serious harms that racism and other forms of discrimination inflict on our communities. That means taking a hard look at our data systems and asking who the data we collect elevates, who is being centered in our data, who is being excluded, and why.

The Commission, a diverse group of 16 members with expertise in areas including healthcare, advocacy, government, business, and public health asked those questions and more. Together, these leaders considered a broad range of actionable solutions for strengthening our public health systems to embrace innovation, inclusion, and anti-racism. The recommendations set forth in this report amount to a blueprint for building a robust 21st century public health data system that advances health equity and equips us to address the racism embedded in our current infrastructure.

It's impossible to fix what isn't measured. We have an opportunity now to create a data infrastructure that is centered on equity and that creates fair and just opportunities for everyone. Let's take that opportunity to promote health and well-being for everyone in America.

RICHARD E. BESSER, MD
President and CEO
Robert Wood Johnson Foundation
### Summary of Recommendations

#### Center health equity and well-being in narrative change
- Identify the stakeholders in public health data system transformation and how to engage them at each step of the transformation process.
- Build on stakeholder identification and develop a campaign to promote the importance of public health data and the need for a transformed public health data system.
- Develop a competencies framework to increase data literacy for various stakeholders about the importance of equity considerations in data systems.
- Build the public health data system needed to shift the narrative to one that is just, positively oriented, and equity-based (e.g., from deficit to strengths, from oppressive to restorative).

#### Prioritize equitable governance and community engagement
- Prioritize and accelerate implementation of the Evidence Act (Foundations for Evidence-Based Policymaking Act of 2018) for improved transparency, quality, and availability of data.
- Establish and implement a coordinated state and federal investment strategy that includes regular fiscal support of state infrastructure coupled with intermediate and long-term system development and data collection.
- Generate and sustain system transformation with defined governance and stewardship models and structures.
- Make sharing and pooling data (at both the individual level and system level) the default for agencies receiving public money and provide data to all actors.
- Build efficient and interoperable data systems to generate comprehensive, complete, and timely data. Collect data with adequate granularity across population groups (inclusive of race/ethnicity, language ability, disability) and geographic levels that are useful at the community level and can be aggregated and disaggregated.
- Develop agile, analytical methods to work with existing data sets and across diverse sets of quantitative/ qualitative data, including historical data.
- Technology companies should support public health data system transformation in under-resourced areas of the country with the largest health inequities, either by direct financial support (corporate social responsibility-CSR) or through skills-based volunteer approaches.
- Philanthropy should fund gaps in public health data, particularly for communities with less resources.

#### Ensure public health measurement captures and addresses structural racism and other inequities
- Build on the Executive Order (EO) on Advancing Racial Equity and Support for Underserved Communities Through the Federal Government: Equitable Data Working Group, to establish an Interagency Data Council, with responsibility for equity, racial justice, & social and public health data.
- As part of public health data system redesign, collect self-reported data by race, ethnicity, income, education, gender identity, sexual orientation, disability, and social position (i.e., how people are placed in a hierarchy of value by society, as perceived by the individual).
- Invest in community relevant and nationally significant metrics on factors that influence health outcomes.
- Collect data that are more accurate and relevant at the community level to enable small area estimates that enable communities and local health departments to prioritize and address local health challenges and measure progress towards healthier communities.
- Develop methods for interpreting public health data that are inclusive of community input, paying attention to messaging, communication, and narrative.
CALL TO ACTION SUMMARY

Government as well as private and other civil society sectors have a role to play in transforming our public health data system.

<table>
<thead>
<tr>
<th>SECTOR</th>
<th>ACTION</th>
</tr>
</thead>
</table>
| **FEDERAL GOVERNMENT** | - Develop minimum standards about data collection, disaggregation, presentation, and access, in federally funded data collection efforts, with an orientation to “freeing federal data” or promoting greater access.  
- Strengthen public health data infrastructure and incentivize the use of new data collection and analytic approaches.  
- Federal funding for data infrastructure should be prioritized to systems that are standards-based and interoperable.  
- Offer guidance on interpretation of racial/ethnic variation in health-relevant data to counter longstanding acceptance of the idea that these variations reflect biological differences rather than systemic, cultural, behavioral, and social factors. |
| **STATE GOVERNMENT** | - Ensure that state policies for public health data collection, sharing (including balancing privacy and transparency), and analysis are equity driven and explicitly call out the influence and consequences of structural racism and other inequities on health.  
- Partner with local health departments and departments that provide public health data (e.g., social services data) to consider new models of collaboration to improve efficiency and timeliness of decision-making and action at both state and local levels. |
| **LOCAL GOVERNMENT** | - Regularly connect public health data to local communications (i.e., what is happening in the community and how it relates to overall community well-being, or the impact of inequity).  
- Ensure the local voice is represented not only in public health data (e.g., from whom data are collected), but in positions of authority responsible for making sense of the data and informing decisions.  
- Explore data-sharing collaborations across government and civil society (e.g., nonprofits, businesses) that can more consistently generate public health data to support equity considerations and advance innovative public-private collaborations on data and analysis. |
| **PUBLIC HEALTH** | - Lead multisector collaboration around public health data sharing to improve the timeliness and quality of data to strengthen local decision-making.  
- Strengthen capacity, diversity, and ongoing training of the public health workforce to monitor and address health equity, both in the field of public health and through novel collaborations with business, academia, or other sectors that influence health.  
- Advocate for and prioritize modernization efforts and data sharing within and across the public health system to ensure that local data can inform emerging public health concerns at the regional, state, and national levels in real time. |
### BUSINESSES
- Work with government partners and other organizations to develop standards through which public health data generated by the private sector can be used and communicated.
- Engage in more consistent workforce exchanges and data partnerships with public health organizations to cross-pollinate innovations in the types, content, quality, and precision of public health data.
- Foster public health innovation by promoting methods in use by technology companies, such as data integration, “big data” analytics, and data security.

### HEALTHCARE SYSTEMS
- Collect social determinants of health data at every consumer encounter, using standardized questions and ICD-10 codes that allow data aggregation within communities.
- Overcome historic silos and build partnerships and legal solutions to facilitate sharing of relevant healthcare data with public health departments in a timely and efficient manner, allowing data linkages and disaggregation of subgroups and geographic regions.

### NONPROFITS
- Build trust in data as a public good in the community and among constituents and ensure that such data do not further marginalize or bring harm to populations.
- Work locally to develop data-sharing strategies with government, and, where appropriate and where there is health benefit, reciprocity, and equitable access.

### PROFESSIONAL ASSOCIATIONS
- Develop strategies to improve data completeness and quality, particularly with respect to equity and drivers of health and well-being.
- Actively work with professional associations from other sectors that influence health (e.g., education, social services, environment) to align efforts, particularly around standardization and data use.

### SCHOOLS
- Help students monitor and evaluate structural and systemic inequities and understand what data systems must include to signal those inequities.
- Train the next generation of public health practitioners and researchers to meaningfully partner with local stakeholders, and to value lived experience and community expertise as much as formal training.

### PHILANTHROPY
- Promote a national, unifying health goal that can be understood and executed, with a focus on the role of structural racism and the values of health equity and well-being.
- Leverage convening and funding opportunities to meaningfully engage leaders, including those from non-health sectors and community organizations, who can advance the importance of public health data.
- Focus funding on key parts of data systems modernization, such as improving governance and using new forms of data and analytic methods to convey better stories of health equity and well-being.
Public health data and data systems are essential to achieving health and racial justice.

Public health data and data systems take the pulse of our society; they measure and indicate how well we sustain the life and vitality of our population and our democracy. As such, these data and systems include governmental public health data and data from many other sectors (e.g., environment, criminal justice, human services, employment, education) and levels (e.g., community, state, nation) that can inform efforts to improve health equity and well-being. Data are the building blocks of population health narratives—stories that emerge from data—that help the nation contextualize what drives or impedes health. What data are collected and how data are collected and interpreted shape these narratives. Data help us identify and understand the magnitude of and reasons for inequity in outcomes and the influence of structural factors like racism and other forms of discrimination on the ability to live a healthy life. To be meaningful, data must reflect accurate and timely information about all population groups and their individual and collective capacities to experience health and well-being.

The year 2020 revealed significant flaws—indeed, fault lines—within our public health data systems’ capacity and infrastructure. COVID-19 exposed these problems explicitly as lack of transparency, conflicting information, and inconsistent and often missing data sowed public distrust, confusion, uncertainty, and fear about individuals’ and the nation’s current and future health.

Now is the time for a reset. Inequities in exposure to COVID-19, illness, and death accelerated awareness of the need to address the disparate health opportunities for diverse racial, ethnic, and other marginalized populations, including immigrants, people with disabilities, returning citizens, and the LGBTQ+ community. This developing awareness was amplified by our collective witness to the brutal murder of George Floyd, an unarmed Black man, by a White police officer. Mr. Floyd’s murder was not an isolated event. However, it catalyzed nationwide recognition that the relentless killings of unarmed persons of color by police and the racial inequities in COVID-19 infection, illness, and death were both manifestations of structural racism—or laws, policies, cultural representations, and norms across interconnected systems that support the unfair treatment of some groups based on the social construct of race that is grounded in a false ideology of a hierarchy of human value.

An unprecedented level of public protest that united people of all races in calling for racial reckoning and justice gives the United States an opportunity to fundamentally change public narratives and institutional policies. Three co-occurring crises (COVID-19, structural racism, and economic disruption for millions of Americans with low incomes) all signal the need for data that can shed light on the structure and intersectional dynamics involved in improving health and well-being and help reframe the national narrative to one that is just, positively oriented, and focused on equity (e.g., from deficit to strength, from oppressive to restorative) from one that is deficit focused.
At the same time, we continue to face many emerging new threats. For example, life expectancy in the United States departed from the trajectory of other wealthy nations decades ago and has dropped in recent years. COVID-19 has fueled that decline. This drop has disproportionately affected people of color, reversing decades of progress in reducing the racial mortality gap.¹

In our current data system, data on health inequities are decontextualized from history and the experiences of race, intersectionality, and place that impact health. Although current systems report on health status and disparities, data can perpetuate health inequities when they lack appropriate context for understanding the root causes of those inequities. We need compelling, inclusive information to help drive critically needed actions and policies. The challenges before us are:

The National Commission to Transform Public Health Data Systems addressed these issues and challenges during its recent deliberations. Convened by the Robert Wood Johnson Foundation (RWJF), the 16-member Commission was charged with reimagining how data are collected, shared, and used, and identifying the public- and private-sector investments needed to modernize our public health data infrastructure and improve health equity. This report summarizes the recommendations that emerged from the Commission’s deliberations. The report is organized in the following sections.

**PART 1** of the report presents the Commission’s recommendations, organized into three major themes. The first theme, **Health Equity and Well-Being Narrative Change**, includes recommendations that emphasize the importance of centering public health as the nation’s pathway to better health and well-being. These recommendations focus on building support among a broad array of stakeholders for a new public health data system that advances health equity, creating the messaging needed to promote the role of data in this process, preparing stakeholders and the workforce to achieve this change, and beginning to create new systems and infrastructure. The second theme, **Equitable Governance, Systems, and Community Engagement**, focuses on the data stewardship, technology, and partnerships needed to transform the system. This section addresses the regulatory, technical, and collaborative work that must be done to create a data system with the breadth, capacity, and specificity needed to understand what drives health and to promote evidence-based decision-making. The **Measuring and Addressing the Health Impact of Structural Racism and Other Inequities** theme addresses the changes needed to redress contemporary and historic health effects of racism and other systemic barriers to opportunity for diverse groups. This section focuses on relationship-building and the types of data needed to transform the system.
Part 1 also includes an **Introduction** that describes the urgency of these changes, the current landscape, and the opportunities for transformation.

**PART 2** of the report describes the steps taken and knowledge gained in preparation for the Commission process. RWJF commissioned the RAND Corporation to produce a series of white papers that anchored this process by describing the current landscape, emerging innovations, and the opportunities for transforming and transitioning to a new public health data system. The papers were informed by an environmental literature scan, review of reports on public health data transformation, and stakeholder interviews. In addition, the papers included insights from 20 organizations that received RWJF grants for innovative work related to data equity and data systems integration. The formative work also included focused discussions with five expert panels on population-specific data gaps (American Indians/Alaska Natives, Blacks/African Americans, LGBTQ+ communities, people living with disabilities, and women). The section goes on to describe the formation of the Commission, the adaptation of the Truth, Racial Healing, and Transformation (TRHT) framework used during the Commission’s deliberations, and the Commission’s work to create a new vision for transforming the public health data system and its recommendations for achieving that vision.

**PART 3** presents a call to action for various sectors and stakeholders, including federal, state, and local governments, and businesses, health systems, nonprofits, schools, associations, and philanthropy. This section provides a preliminary blueprint of when, how, and by whom actions can be taken to implement and advance and build on these recommendations.

---

**What is a modern, transformed public health data system?**

The Commission defines a transformed public health data system as one that is accountable to and reflects the perspectives of local communities and diverse populations, including people of color, immigrants, persons with disabilities, and the LGBTQ+ community. The system must be sustainable, fully integrated with healthcare and other sectors that drive health by linking data across multiple sectors and at multiple levels, and provide data that are disaggregated, comprehensive, and timely. It should move from problem-focused to solution-focused analysis and have the capacity to provide the knowledge needed to optimize health and well-being for all people. It must provide tools to address racism and racial/ethnic and other social disparities. It should facilitate restorative systems that respect all communities, including the sovereign right of tribal nations to govern data and ensure they have access to it; protect individuals’ privacy and security; and guard against unintended consequences.
Members of the independent National Commission to Transform Public Health Data Systems

**GAIL C. CHRISTOPHER,**
(Commission Director), DN, Executive Director, National Collaborative for Health Equity

**MARGARITA ALEGRIA,**
PhD, Chief of the Disparities Research Unit, Massachusetts General Hospital; Professor, Department of Psychiatry, Harvard Medical School

**MARY T. BASSETT,**

**RAYMOND BAXTER,**
PhD, Co-Chair, Roundtable on Population Health, National Academies of Sciences, Engineering, and Medicine

**JULIET K. CHOI,**
JD, President and Chief Executive Officer, Asian & Pacific Islander American Health Forum (APIAHF)

**MICHAEL CRAWFORD,**
MBA, MHL, Associate Dean for Strategy, Outreach, and Innovation (ADSOI), College of Medicine, Howard University

**FERNANDO DE MAIO,**
PhD, Director, Research and Data Use, Center for Health Equity, American Medical Association, and Professor of Sociology, DePaul University

**KAREN DESALVO,**
MD, MPH, MSc, Chief Health Officer, Google

**ABIGAIL ECHO-HAWK,**
MA, Director, Urban Indian Health Institute; Chief Research Officer, Seattle Indian Health Board

**THOMAS LAVEIST,**
PhD, Dean, School of Public Health and Tropical Medicine, Tulane University

**ALEXIS C. MADRIGAL,**

**JOHN LUMPKIN,**
MD, MPH, President, Blue Cross and Blue Shield of North Carolina Foundation

**AMY O’HARA,**
PhD, MA, Research Professor, Massive Data Institute, and Executive Director, Federal Statistical Research Data Center, McCourt School for Public Policy, Georgetown University

**JONATHAN PERLIN,**
MD, PhD, Chief Medical Officer and President, Clinical Services, HCA Healthcare

**NINEZ PONCE,**
MPP, PhD, Director, UCLA (University of California Los Angeles) Center for Health Policy Research; Principal Investigator, California Health Interview Survey; Professor, Department of Health Policy and Management, UCLA Fielding School of Public Health

**CHESLEY RICHARDS,**
MD, MPH, Retired Deputy Director for Public Health Science and Surveillance (DDPHSS), Centers for Disease Control and Prevention (CDC)

**JAVIER ROBLES,**
JD, Faculty Member and Professor, Kinesiology and Health Department, and Director, Center for Disability Sports, Health and Wellness, Rutgers University
RECOMMENDATIONS
1. Center Health Equity and Well-Being\textsuperscript{a} in Narrative Change

The approach to collecting, analyzing, and using public health data must be motivated by a new narrative that prioritizes health equity and factors in all of the physical, social, mental, cultural, economic, and environmental drivers of health and well-being. A modern and transformed public health data system would address structural racism, equity, and well-being considerations as the guiding influence in all data-informed decisions.

Recommendation 1a. Identify the stakeholders in public health data system transformation and how to engage them at each step of the transformation process.

\textbf{CONTEXT.} A modern, equity-oriented public health data system should mobilize the full range of actors and sectors that influence health outcomes, help identify health priorities, and catalyze action. There is growing evidence about the benefits of having full community and multisectoral participation in public health decision-making.\textsuperscript{3} But too often groups that have been marginalized and disproportionately affected by health inequities are excluded.\textsuperscript{4} In the context of new forms of data or “big data,” issues of participation and representation are even more critical.\textsuperscript{5} Best practices for community engagement must be used to facilitate authentic engagement and empower community members and stakeholders.

\begin{itemize}
  \item Government leaders
  \item Tribal leaders
  \item Health care systems
  \item Private sector/Business leaders
  \item Media leaders
  \item Philanthropies
  \item Community-based organizations
  \item Academia/Research institutions
  \item Advocates for diverse communities and population groups
\end{itemize}

\textsuperscript{a} “Well-being means thriving in every aspect of life and having opportunities to create meaningful futures. It includes people’s physical, mental, and social health, as well as basic needs like food, housing, education, employment, and income. It includes social and emotional needs, like sense of purpose, safety, belonging and social connection, and life satisfaction. It must include freedom from racial oppression, and active advancement of racial equity. The concept is tightly linked with the well-being of our communities, our environment, and our planet. While it often has been incorrectly considered an individual wellness concept, well-being goes far beyond that by considering relationships among people, interconnectivity of cultures and societies, and relationships between people and the planet.” From: Setting a new narrative about progress and well-being, RAND and Metropolitan Group, August 2021.
Recommendation 1b. Build on stakeholder identification and develop a campaign to promote the importance of public health data and the need for a transformed public health data system.

**CONTEXT.** Much of what are currently considered public health data is stored by local public health departments, healthcare organizations, and other entities outside health that generate social determinants of health (SDOH) information. These data are not fully available or accessible for the diverse stakeholders comprising the broader public health system. Thus, the value of public health data is not always well understood. Further, many sources of public health data fall outside of health (e.g., departments and organizations related to social services, economics, environment). A public health data system should ensure open access where appropriate, provide supporting materials to support meaningful engagement with data, link to actionable policy and programmatic solution options, be designed to invite access and usability, and be useful to people working to improve health in their communities.

**WHAT NEEDS TO BE DONE**

**SHORT-TERM**
- Develop case studies that explain the story and the value of public health data. Potential case studies include the 2020 Census, which showcases how data drive resource allocation, and COVID-19, which can illuminate the role of data in tracking a global pandemic.
- Develop messaging aimed at different audiences about the value of public health data and modernizing the public health data system. Use microtargeting, representative storytelling, and economic framing (meaning tighter links between public health, economics, and well-being) to carry out messaging and subsequent campaigns.
- Identify diverse leaders (e.g., from professional associations, grassroots organizations), across all U.S. geographies and subgroups, who will serve as campaign ambassadors and prepare them for that role.

**LONG-TERM**
- Highlight the important role of collecting and analyzing public health data for implementing equitable public policy.
- Continue to emphasize the value of public health data and its linkage with community and national progress.

**WHO NEEDS TO ACT**
- Government leaders
- Tribal leaders
- Private sector/Business leaders
- Media leaders
- Professional associations (e.g., National Association of County and City Health Officials, American Hospital Association, National Association of Community Health Centers and American Essential Hospitals, American Medical Association)
- Philanthropies, such as RWJF
- Academia/Research institutions
- Grassroots and community leaders
Recommendation 1c. Develop a competencies framework to increase data literacy for various stakeholders about the importance of equity considerations in data systems. This framework also can deepen understanding of the value of public health data, shift perceptions about what constitutes public health data and who has access to them, and explore how data can motivate public health data system transformation.

**CONTEXT.** There is growing concern that the public health workforce lacks the core skills and competencies to procure and optimize information technology. Individuals skilled in informatics are choosing private industry over public health for better pay and advancement. The public health workforce could benefit from expanding analytic competencies through a multipronged approach (e.g., school course offerings, narratives about innovation in public health data, and workforce exchanges with the private sector) to build competencies and fill this gap. Several federal policies (e.g., 2020 Federal Data Strategy) can be leveraged for this workforce support.

“Misinformation is a serious issue. It is important we create an environment where all communities trust data.”
– Commissioner Michael Crawford

**WHAT NEEDS TO BE DONE**

**SHORT-TERM**
- Train the public health data workforce to understand why equity should be embedded in the content, structure, analysis, translation, presentation, access, use, and dissemination of data.
- Make sure the workforce embraces diverse perspectives on how to shift data and their accompanying narratives toward health equity and well-being.

**LONG-TERM**
- Permanently shift the public health narrative by integrating the value of public health data and their role in social change into the curriculum of graduate/undergraduate schools (e.g., public health, social work, health professions, public policy, economics, sociology, political science, business, environmental science) and other training programs.

**WHO NEEDS TO ACT**
- Federal government leaders
- Public health departments
- Health care systems
- Private sector/Business leaders, particularly from technology and data generation companies
- Professional associations
- Philanthropies
- Academia/Research institutions
Recommendation 1d. Build the public health data system needed to shift the narrative to one that is just, positively oriented, and equity-based (e.g., from deficit to strengths, from oppressive to restorative).

**CONTEXT.** The current public health data system disproportionately focuses on negative health outcomes (e.g., mortality and morbidity), and overlooks positive health and well-being measures (e.g., community cohesion, hope, civic engagement, prosocial behaviors, resilience). Without such information, the nation cannot track aspects of thriving and flourishing that are central to health and well-being. A future public health data system also must monitor the processes and outcomes of inequity, history, cumulative risk, and cumulative trauma on health. The modern public health data system must look both upstream, at the structural drivers of health inequities, and over time, at intergenerational inequities.

**WHAT NEEDS TO BE DONE**

**SHORT-TERM**

- Assess gaps in the current data system (e.g., standard race/ethnicity reporting at different levels of government and across agencies, user interface, security and privacy, usability for different languages and literacies), and how these gaps may impede a more holistic and equity-centered public health narrative.

- Identify minimal datasets to tell narratives of equity that are specific to the needs and context of diverse communities (e.g., immigrants, individuals who are incarcerated, people released from prison, people not actively seeking employment or housing or healthcare, people with disabilities) as well as to address issues of race/ethnicity, disability, language access, and American Sign Language (ASL) needs.

**LONG-TERM**

- Create specific protocols and multiple use cases for a transformed public health data system that supports restorative and prospective storytelling. Data should be complete enough to showcase history and context, how communities respond to challenges, and should feature community voices.

- Use those cases to inform public health data narrative change strategies. For example, how accurate representations of diverse communities and the effects of the COVID-19 pandemic are utilized to foster timely and appropriate responses to needs.

- Ensure that efforts to increase public health infrastructure capacity prioritize support for building an effective public health data narrative focused on health equity and well-being.

**WHO NEEDS TO ACT**

- Office of the National Coordinator for Health Information Technology (ONC), Department of Health and Human Services (HHS)

- White House, Data Equity Working Group and Office of Management and Budget (OMB)

- Private sector/Business leaders, particularly from technology companies

- Hospital associations

- Philanthropies

- Academia/Research institutions

- Other members assembled through a Federal Advisory Committee process

---

b. A use case is a description of all the ways an end-user wants to “use” a system.

c. Restorative storytelling is an approach to storytelling in which descriptions show how people and communities respond and recover after difficult times; these stories do not just focus on challenges.
2. Prioritize Equitable Governance and Community Engagement

A modern public health data system must execute a plan for governance, decision-making, and community engagement that centers addressing structural racism and creating equity in its design and operation. This means transparency, accessibility, and interoperability in all aspects of how the public health data system runs, how the system interacts with other systems, and how it takes in new information to respond to emerging and ongoing health issues.

**Recommendation 2a.** Prioritize and accelerate implementation of the Evidence Act (Foundations for Evidence-Based Policymaking Act of 2018) for improved transparency, quality, and availability of data.

**CONTEXT.** The Foundations for Evidence-Based Policymaking Act (Public Law No: 115-435)\(^{10}\) requires agency data to be accessible and requires agencies to plan to develop statistical evidence to support policymaking. It also includes the OPEN Government Data Act, which “requires public government data assets to be published as machine-readable data” and an online federal data catalogue and “a single point of entry for the public to access agency data.”\(^{11}\) (These laws do not apply to tribal nations.) Accelerating the law’s implementation would provide federal momentum for an equity-oriented data system. Other federal activities to address equity and upstream drivers of health could be further leveraged to strengthen the public health data system. Examples include:

- **EXECUTIVE ORDER (EO) ON ADVANCING RACIAL EQUITY AND SUPPORT FOR UNDERSERVED COMMUNITIES.** This EO calls for data to be disaggregated by race, ethnicity, gender, disability, income, veteran status, or other key demographic variables to advance equity.\(^{12}\)

- **INTERDEPARTMENTAL HEALTH EQUITY COLLABORATIVE (IHEC) DATA WORKGROUP.** The IHEC Data Workgroup includes experts engaged in data activities and data policies to address SDOH and health disparities. The goals of the group are to identify existing policies and practices for improving access to data and use of data in support of policy development and to promote data projects and applications that address SDOH and support efforts to advance health equity.\(^{13}\)

**WHAT NEEDS TO BE DONE**

**SHORT-TERM**
- Integrate Commission recommendations with the above efforts, leveraging Commissioner connections and positions on various workgroups.

**LONG-TERM**
- Advocate for and support federal, state, and local policies that promote transparency, quality, and availability of data in a manner that also respects indigenous data sovereignty, which is the right of a nation to govern the collection, ownership, and application of its own data. It derives from tribes’ inherent rights to govern their people, lands, and resources.\(^d\)

**WHO NEEDS TO ACT**
- Federal government leaders
- Tribal leaders
- Academia/Research institutions
- Transforming Public Health Data Commissioners, in supporting role

\(^d\) [https://usindigenousdata.org/]
Recommendation 2b. Establish and implement a coordinated federal, state, and local investment strategy that includes regular fiscal support of state infrastructure coupled with intermediate and long-term system development and data collection.

**CONTEXT.** General federal and state funding for public health has declined over the past decade. The resulting wide variability in public health infrastructure has had critical cascading effects for health equity. Key to achieving an equitable health data system is assessing how to migrate legacy data systems to efficiently participate in any data sharing model with minimal human effort and securing the funding and other supports needed to sustain these new systems. For example, COVID-19 has driven the rapid adoption of electronic case reporting, which could immediately add data about more than 100 notifiable and reportable conditions. However, state agencies with multiple legacy surveillance systems often cannot receive and parse this real-time reporting. Although federal policy and funding can shape a new vision of a modern, equity-oriented data system, state governments can reinforce the value of equity orientation in data or counter them through restrictive policies and narrow resource allocation decisions.

**WHAT NEEDS TO BE DONE**

**SHORT-TERM**
- Secure short-term investment in infrastructure, including technology and necessary human capabilities, to leverage data for decision-making.

**LONG-TERM**
- Secure longer-term investments to migrate legacy data systems and ensure that systems are sustained and maintained.

**WHO NEEDS TO ACT**
- Federal government leaders
- State government leaders
- Private sector/Business leaders
Recommendation 2c. Generate and sustain system transformation with defined governance and stewardship models and structures. These should sustain and hold accountable a transformed data system that supports equitable and just narratives and inclusive decision-making.

**CONTEXT.** The rapid pace of innovation in information technology and changes in what, how, when, and why data are collected, and by whom, raise important questions. What governance structure is the best guide for our current public health data system? How can we ensure that new governance structures are nimble enough to address future health data considerations? A strong governance structure could prevent inappropriate use of public health data, when individual users are not good stewards of the resources to which they have access. In addition, concerns that state and local preemption of federal sources of public health data with other data (or vice versa) has created misalignment and limited the acknowledgement of health and economic inequities. This tension has elevated scrutiny of the relationship between state and local governments and governance models.

<table>
<thead>
<tr>
<th>WHAT NEEDS TO BE DONE</th>
<th>WHO NEEDS TO ACT</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SHORT-TERM</strong></td>
<td>Federal government</td>
</tr>
<tr>
<td>- The CDC should get input from states/localities/tribes and align goals and funding to ensure that governance and improved infrastructure robustly support local goals and needs.</td>
<td>CDC</td>
</tr>
<tr>
<td>- Under executive order, appoint a chief health strategist for the country.</td>
<td>Federal, state, local, tribal, and territorial public health leaders</td>
</tr>
<tr>
<td>- Establish a higher-level body that could serve advocacy and oversight functions. This body could also oversee implementation of recommendations from the Commission and other related parties.</td>
<td>Private sector/Business leaders</td>
</tr>
<tr>
<td>- Set up advisory groups at multiple levels and with equitable representation to provide input on and establish a governance structure. Key decisions should seek to achieve a more equitable balance in governance, ensure power is given to communities (including identity-based groups), and ascertain what governance looks like at each level (federal, state, local, tribal, territorial).</td>
<td>Philanthropies</td>
</tr>
<tr>
<td><strong>LONG-TERM</strong></td>
<td>Communities/community members</td>
</tr>
<tr>
<td>- Develop enforcement mechanisms to ensure sustained engagement around the principles of a transformed public health system.</td>
<td></td>
</tr>
<tr>
<td>- Enact statutory requirements for supporting equity in all policies.</td>
<td></td>
</tr>
<tr>
<td>- Generate long-term commitment from funders/investors.</td>
<td></td>
</tr>
</tbody>
</table>
Recommendation 2d. Make sharing and pooling data (at both the individual level and system level) the default for agencies receiving public money — because no one sector or institution holds all the data needed to understand the factors that drive inequities in health and well-being — and provide data to all actors.

**CONTEXT.** Equity in data access must ensure that the system is structured equitably so that everyone has the same level of access to the same information. Tying data expectations to federal payments is a well-established practice that could be expanded. In recent years, for example, the federal government has leveraged incentive programs to promote interoperability and the collection of a standardized set of data through the Centers for Medicare and Medicaid Services’ (CMS) Meaningful Use program and, more recently, the Merit-based Incentive Payment System (MIPS). These incentive programs have tied provider payments to standards of data capture and information exchange. In 2020, the Office of the National Coordinator for Health Information Technology (ONC) released the CURES Act Final Rule “designed to drive interoperability of EHI by supporting the use of ... Fast Healthcare Interoperability Resources (FHIR) standards for application programming interfaces (APIs).” Use of FHIR, and recent federal laws, executive orders, and other federal activities fosters data sharing among a wide range of potential users, including patients, providers, and other health care entities.

> “The transformed system has to embody the ability to link data across multiple sectors and at multiple levels, from the planet to the neighborhood in order for us to understand how we got here, where we are right now, and what we are facing ahead.”
> 
> – Commissioner Raymond Baxter

<table>
<thead>
<tr>
<th>WHAT NEEDS TO BE DONE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SHORT-TERM</strong></td>
</tr>
<tr>
<td>▪ Enable linkage of data from multiple cross-sector sources with governance oversight (e.g., privacy, security, and anti-discrimination protections) that includes representatives from groups most at risk for harm from data misuse.</td>
</tr>
<tr>
<td>▪ Governance of data sharing should be driven at the local/state/tribal level, where the laws and the needs of the population vary dramatically, and should inform what information can be shared, with whom, and under what conditions.</td>
</tr>
<tr>
<td>▪ Address proprietary and other disincentives to sharing data and implement public accountability mechanisms functions where needed.</td>
</tr>
<tr>
<td>▪ Implement the Information Blocking Rule, which prohibits practices by healthcare providers and others that are likely to interfere with, prevent, or discourage data access, exchange, or use of electronic health information.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>WHO NEEDS TO ACT</th>
</tr>
</thead>
<tbody>
<tr>
<td>▪ Government leaders</td>
</tr>
<tr>
<td>▪ State and local public health officials</td>
</tr>
<tr>
<td>▪ Private sector/Business leaders</td>
</tr>
<tr>
<td>▪ Academia/Research Institutions</td>
</tr>
<tr>
<td>▪ Communities/community members</td>
</tr>
<tr>
<td>▪ Commission members and other experts</td>
</tr>
</tbody>
</table>
WHAT NEEDS TO BE DONE (CONTINUED)

- Develop a federal mechanism to create interoperability standards for public health, community health, and healthcare.
- Develop a consensus statement on definitions and priorities for sharing data on measures outside public health (e.g., food security, violence, criminal justice).
- Build capacity for shared services so that communities can benefit from advances in data pooling, integration, visualization, and GIS, and build trust in sharing data.
- Involve stakeholders with appropriate expertise in training and coaching the public health workforce and supporting small public health departments through internships and skills-based volunteering to address staffing shortfalls.

LONG-TERM

- Develop a governance mechanism in consensus with relevant stakeholders and communities that addresses concerns about trust, privacy, confidentiality, and security, strengthens confidence in the process, and provides transparent choice and control.
- CMS should make conditions of participation in Medicare and Medicaid contingent on sharing data and should create a national Medicaid data set.
- Propose policy recommendations to authorize tax or other incentives for companies to offset costs and perceived disadvantages of sharing anonymized data and provide support for best practices in anonymizing data.
- Create a structure and incentives for health systems to collect critical SDOH data that follows the same equity principles suggested for government data.

“We need metrics that give us a pulse on what is happening in the community, as opposed to a rearview mirror look.”

– Commissioner Karen DeSalvo
Recommendation 2e. Build efficient and interoperable data systems with guidance on standardized data collection and rollup of granular categories, if needed, to generate comprehensive, complete, and timely data. Collect data with adequate granularity across population groups (inclusive of race/ethnicity, language ability, disability) and geographic levels that are useful at the community level and can be aggregated and disaggregated. Disaggregation should include multiple factors by race, ethnicity, disability, gender, and others.

CONTEXT. Public health recommendations are often made because of emerging needs, based on evolving, incomplete, and sometimes imprecise public health data. However, timely and precise (defined as exact and/or sharply defined) data are scarce, in large part because of the resources required. Ensuring an equity-oriented data system requires careful consideration about what types of data are collected and at what level of granularity, and how to balance valid concerns around individual privacy with the benefit of disaggregated data to inform public health and policy solutions. This point is particularly salient for some populations, which may be at risk if such data were made publicly available (e.g., undocumented residents, individuals with disabilities). The lack of data granularity poses challenges for understanding what works, when, and for whom, but there is no consensus on the level of geographical and other data granularity needed to support public health data decisions at the national versus local level.

WHAT NEEDS TO BE DONE

SHORT-TERM

- Convene the public and private sectors to develop a national strategic plan for data modernization, including priorities, necessary data streams, and data governance, that cuts across sectors and examines existing structures and systems.
- Investigate longitudinal health records, “data lakes” (systems or repositories of data stored in their natural or raw format), and synthetic data to refine algorithms and advance small-area estimation and the need for an opt-in or consent process for individual health records.

LONG-TERM

- Build connections to global data systems and address governance issues. Using the Health Information Technology for Economic and Clinical Health Act (HITECH) model, the CDC should contract with academic institutions through rapid grants to develop synthetic data lakes.
- Explore issues with consent for data donation to public health entities (e.g., health information exchanges, all-payer claims databases).

WHO NEEDS TO ACT

- The administration, led by the White House’s Office of Science and Technology (OSTP) in collaboration with CDC and ONC
- HHS Office for Civil Rights
- Department of Commerce
- Census Bureau
- Federal Communications Commission (FCC)
- Academia/Research institutions
Recommendation 2f. Develop agile, analytical methods to work with existing data sets and across diverse sets of quantitative and qualitative data, including integrating historical data and addressing any issues of algorithmic and related bias.

**CONTEXT.** Data integration across a wide range of sources, the sheer volume of health-related data, and increased computing power and technological innovation together hold great promise for the development of proactive, data-driven solutions that improve health, equity, and well-being. With these changes, however, comes a need for new methodologies to analyze data efficiently, cost-effectively, and accurately. To ensure that data governance and privacy protections keep pace with information technology innovation, methodologic advancements tied to tiered access and data de-identification that reduce the likelihood of re-identification could also be explored. Methodologic approaches that allow disaggregation and analysis of data by geography or population characteristics could also help advance health equity.

**WHAT NEEDS TO BE DONE**

**SHORT-TERM**

- The CDC should continue to work with researchers and technology leaders to reimagine a broader methodological approach to analyzing and leveraging health data that incorporates novel methods, signals, etc., through the lens of structural and social determinants of health and health equity. This effort should focus on improving local and smaller-area data.
- Devote resources to devise better methods to pool and use existing data.
- Identify and address the bias and harm of current methodological paradigms by moving from problem-focused to solution-focused analysis. This requires guidance and protocols for understanding and using race and ethnicity as variables in health research and doing more to support research that measures the health effects of racism.
- Designers of algorithms and place-based predictive metrics should assess the impact of those algorithms across racial/ethnic groups, and whether these metrics provide useful specificity and sensitivity and/or have limitations.
- Use novel measures that detect emergent trends or conditions, such as those surfacing on contemporary technology platforms, to capture sentiment and to track any public concerns about disinformation or misuse of data.

**LONG-TERM**

- Develop a code of ethics for novel measures work and other advanced methodologies and technologies, to ensure appropriate interpretation and use of the data.

**WHO NEEDS TO ACT**

- CDC
- Philanthropies and government to fund methodological advances
- Research methodologists (e.g., National Academies of Science, Engineering, and Medicine) and data scientists

---

Recommendation 2g. Technology companies should support public health data system transformation in under-resourced areas of the country with the largest disparities in opportunities for health and health outcomes, either by direct financial support (corporate social responsibility) or through skills-based volunteer approaches.

**CONTEXT.** Technology companies’ interest in health has continued to grow, with the expansion of smartphones, wearable devices, and application of medical devices (the Internet of Bodies). Very few data from such devices, however, are consistently used for the public good to identify emerging health needs or to inform local decision-making—partly because they tend to be about individuals rather than about populations, geographies, and societal structures and conditions, and partly because technology companies have not committed to public access. Technology companies also have a wealth of talent and are often at the cutting edge of new technologies and approaches to finding signal value (what the data are clearly showing or prioritizing) within vast amounts of data. At the same time, a larger political and societal question focuses on the role of big technology companies like Google, Facebook, Microsoft, and Twitter. The public conversations and philosophical questions about how technology companies should behave and what their role in society should be may open a new door for companies to leverage their data, resources, and expertise for public good, and to become powerful allies in crafting a modern, equity-oriented data system.

**WHAT NEEDS TO BE DONE**

**SHORT-TERM**
- Stakeholders should define roles for engaging the business community.
- Train corporate technology sector boards to improve health equity competencies, engage in discussions, support, and address what is needed for communities and different populations. Incorporate public health data sharing in Environmental, Social, Governance (ESG) standards.
- Expand broadband into rural and urban communities.

**LONG-TERM**
- Delineate responsibilities of technology companies to foster a national public health data strategy.
- Engage and support representation of racial, ethnic, and gender minorities and disability communities to diversify the technology field.
- Build connections with corporate responsibility and data for good programs.
- Develop strategies for under-resourced areas to gain parity with the rest of the country.

**WHO NEEDS TO ACT**
- Regulators
- Technology industry
- Technology stakeholders
- Other private sector/business leaders
Recommendation 2h. Philanthropy should fund gaps in public health data, particularly for communities with fewer resources.

**CONTEXT.** National and regional philanthropic organizations can help catalyze a modern, equity-oriented data system. Although philanthropies cannot, and should not, underwrite the full scope of data modernization efforts, they can invest in methodologic advancements and analysis on critical issues that have blocked equity data system transformation. Through grantmaking, philanthropic organizations can help guide a national or regional agenda by addressing important gaps and needs (e.g., positive health and well-being, structural inequities). Philanthropy can also bring together diverse stakeholders to share ideas, challenge norms, and generate new solutions, and share and support the spread and uptake of evidence-based solutions using public health data.

**WHAT NEEDS TO BE DONE**

**SHORT-TERM**
- Identify gaps in funding.
- Buy access to private data sources for communities and researchers.
- Support convening and consensus around how data on race and ethnicity and other population groups are collected, operationalized, and analyzed.

**LONG-TERM**
- Regional and local philanthropies should partner with national foundations to support the digital transformation of state and local public health systems. This should include improving the capacity for standards-based, bidirectional flow of data and information between public health and healthcare.
- Philanthropy should assist rural and urban providers in historically marginalized, underserved, and under-resourced communities in adopting and participating in the transformed public health data system.

**WHO NEEDS TO ACT**
- Philanthropies
3. Ensure Public Health Measurement Captures and Addresses Structural Racism and Other Inequities

Currently, the public health data system minimally tracks information to guide the dismantling of structural racism and address other inequities that influence health. The modern, transformed public health data system must consistently have data that measure the structural and historical factors driving persistent health inequities and that can be disaggregated and acted upon across sectors and levels of government.

“We need better ways of capturing the concept of structural racism, but most of all we need the field to discipline itself to change the narrative of measured racial inequities to include racism and abandon reflexive pursuit of biological/genetic explanations.”

– Commissioner Mary T. Bassett
Recommendation 3a. Build on the Executive Order (EO) on Advancing Racial Equity and Support for Underserved Communities Through the Federal Government: Equitable Data Working Group\(^f\) to establish an interagency data council, with responsibility for equity, racial justice, and social and public health data. The interagency council would prioritize structural and systemic drivers of health and inequities, establish a process for using public health data, and create a longitudinal and sustainable program that will ensure that resources are available to maintain, improve, and periodically report on the council’s progress.

**CONTEXT.** The COVID-19 pandemic has illuminated the need for an improved public health data system and mobilized action at the local, state, and federal levels. Coalitions and collaboratives focused on interoperability and data sharing have a renewed sense of urgency. Harnessing the collective action and power of these groups to develop a governance structure and path forward toward an equity-oriented public health data system would be a good next step. In 2007, the Agency for Healthcare Research and Quality (AHRQ) put forward the idea of a national health data stewardship entity.\(^{28}\) Although AHRQ did not move forward with the idea, a convening council or other body could engage diverse sectors from social services, education, justice, the environment, along with healthcare and public health.

**WHAT NEEDS TO BE DONE**

**SHORT-TERM**
- Per the EO, establish a sustainable interagency data council, with a charge of improving measures needed to assess racial justice and bring together the different agencies with commitments to creating interoperable public health data.
- Develop standards and expectations around equity, data access, and interoperability for federal public health data.
- Include people from community-based and tribal organizations on the council and related federal agency data committees.
- Advocate for secure and efficient linkages with race and Hispanic origin data through the National Secure Data Service.

**LONG-TERM**
- Examine the role of technology in data interoperability and analysis, with attention to developing guidance for addressing algorithmic bias.
- Propose policies that ensure the autonomy of public health data, including related efforts, such as the Census, by placing it under bipartisan oversight. Public health relies on the Census for key measures of population health, including calculation of birth, death, and fertility rates.

**WHO NEEDS TO ACT**
- Office of the Vice President
- HHS, including CDC
- Office of Management and Budget (OMB)
- Congress

---

Recommendation 3b. As part of public health data system redesign, collect self-reported data by race, ethnicity, income, education, gender identity, sexual orientation, disability, and social position (i.e., how people are placed in a hierarchy of value by society, as perceived by the individual and by others). The data could be used to identify areas of disadvantage where investment and action are needed.

**CONTEXT.** The United States does not consistently or uniformly collect data critical to understanding the health, well-being, or lived experiences of millions of residents, severely limiting our ability to develop data-driven solutions for minority and other vulnerable populations. Even in the midst of a worldwide pandemic, as of August 2020, data were not made publicly available and basic demographic characteristics such as race and ethnicity were not collected or were unknown for 51 percent of COVID-19 cases. These lapses masked severe inequities in case and fatality rates early on and delayed the development and implementation of data-driven solutions. The lack of granularity in data and data measures impedes advancement of population health and well-being. Yet there is no consensus on the level of geographical and other data granularity needed to support public health data decisions at the national and local levels. For example, some have argued for greater granularity of ethnicity classifications to capture sub-group variations in healthcare, risk factors, and health behaviors. Many data on race, ethnicity, language, sexual orientation, and gender identity originate in the healthcare delivery and financing sectors, which face barriers to data collection due to state and federal restrictions.

**WHAT NEEDS TO BE DONE**

**SHORT-TERM**

- Update and clarify the Office of Management and Budget (OMB) Standards for the Classification of Federal Data on Race and Ethnicity (Statistical Policy Directive No. 15). Consider Directive 15 a floor and encourage further data disaggregation within major race and ethnicity categories.
- Update and promulgate the recommendations of the National Academy of Medicine report on standardizing race, ethnicity, and language data.
- Assess whether local, state, and federal laws and regulations impede race, ethnicity, language, sexual orientation, and gender identity data collection and to strengthen privacy protections for those data.
- Encourage the collection of sociodemographic characteristics during all healthcare encounters in ways that preserve trust between providers and consumers.

**LONG-TERM**

- Include measures of structural racism as well as broader positive health metrics, such as in federal and state health-related statistical reports.
- Stratify reporting of hospital quality data by race, ethnicity, disability status, sexual orientation, and gender identity. CMS conditions of participation should address missing race/ethnicity data.

**WHO NEEDS TO ACT**

- National Center for Health Statistics
- CDC
- Office of Minority Health
- CMS
- National Institutes of Health (NIH)
- National Academy of Medicine (NAM)
Recommendation 3c. Invest in community-relevant and nationally significant metrics on structural factors that influence health outcomes, focusing on upstream causes of inequity by measuring how systems segregate, discriminate, and exclude.

CONTEXT. Despite increasing amounts of public health data being collected, data on the upstream factors that influence health are not regularly or consistently available to inform decision-making. A review of the Healthy People 2030 Leading Health Indicators suggests continued gaps in public health data, particularly with regard to economic stability, neighborhood and the built environment, education, and community and social context. Public health data should place more emphasis on social context and other key factors (e.g., education, housing instability, food insecurity) that have a large impact on populations with vulnerabilities and place less emphasis on traditional health data that focus primarily on disease and disease burden.\textsuperscript{33,34} Data on lived experiences and community history (e.g., narratives), finance (e.g., mortgage acceptance rates, redlining, school budgets, county budgets, disability discrimination, language access), and racism and discrimination could provide critical insights into some of the root causes of health inequities.\textsuperscript{35}

“Data systems are often designed to drive interest from above and not for use by local communities.”

– Commissioner Alexis C. Madrigal
**WHAT NEEDS TO BE DONE**

**SHORT-TERM**
- Identify a parsimonious (i.e., minimal and simplest) set of metrics to analyze the health effects of structural racism.
- Develop outcome measures that benchmark a newly defined set of parsimonious lead indicators of racial equity, inclusion, and well-being, and track change over time.

**LONG-TERM**
- Develop new measures that go beyond aggregating individual data to detect structural factors and systemic practices that separate people from health and opportunity. These metrics should provide insight on both assets and deficits that reduce and amplify separation, respectively. This requires methodologic advances to refine metrics for measuring segregation and other forms of separation (including measures of discrimination and structural racism) in institutions and systems, place-based measures that tell the full story (e.g., not masking gentrification), and a clearinghouse for best evidence. Expand frameworks for analyzing individual and community-level data in ways that support valid inferences.
- Measure upstream and downstream influences on public health, using longitudinal data to capture the history of separation and how the forces of segregation have affected people and places. Measure how policies affect systems and places, how these in turn affect people and health, and how each is impacted by interventions. Develop theoretical and empirical models to describe these processes of structural violence, acknowledging the intersectional nature of oppression in U.S society.
- Systematically measure and report racism in all its forms (internalized, interpersonal, institutional, and structural). Aggregate new or currently collected data, including self-reported experience with racism, measurement of discrimination, housing discrimination, and other measures like school exclusionary discipline and gun violence.\(^6\)
- Incorporate these measures into value-based alternative payment models in Medicare, Medicaid, and private insurance plans.
- Create written guidance on interpretation of racial/ethnic health disparities that recognizes “racism as the mechanism by which racial categorizations have biological consequences.”\(^36\)
- Develop guidelines to support improved measurement and analysis of structural drivers of health inequities in community health needs assessments by not-for-profit hospitals and public health departments, as well as community development financial institutions.

**WHO NEEDS TO ACT**
- OMB
- AHRQ
- Congress
- Academia/Research institutions

---

\(^{g}\) Bi-annual AHRQ report: https://www.ahrq.gov/workingforquality/reports/index.html
Recommendation 3d. Collect accurate, relevant community-level data that support small-area estimates so that communities and local health departments can better prioritize and address local health challenges and measure progress toward healthier communities.

Context. The public health data system must have both flexibility and interoperability; it must allow for local flexibility while ensuring that data collected locally can be easily combined with data collected elsewhere. Modularity connotes “the degree to which a system’s components can be separated and recombined and refers to … the degree to which the rules of the system architecture enable or prohibit the mixing or matching of components.” In contrast, systems consolidation or integration is another approach to ensuring interoperability, but it is less flexible in meeting emerging or unique needs. Standardized measures would not only support interoperability but would also allow stakeholders to select those measures most relevant to their work, greatly increasing the flexibility of the public health data system.

“An equity-focused data system will not just describe problems; it will help us work towards solutions. It will be a valuable tool for social justice.”

– Commissioner Fernando De Maio

WHAT NEEDS TO BE DONE

SHORT-TERM

- Guide local leaders in identifying parochial public health measures and data priorities, with consideration of uniformity to support analysis and interoperability over time, including interoperability at intra-local levels, which can be rolled up optimally to the state level.

LONG-TERM

- Provide supports to a local data workforce that can tailor data collection efforts locally and employ small-area estimations and other techniques.
- Ensure that the National Secure Data Service can support small-area estimation needs through access to administrative data that lend strength to local data to produce statistics.

WHO NEEDS TO ACT

- Census Bureau
- National Center for Health Statistics
- NIH
- Department of Housing and Urban Development
- Departments of Education and Public Health
- Department of Agriculture
- Department of Commerce
- Parallel agencies at state and local levels
- Municipal and county planning agencies
- Academia/Research institutions
Recommendation 3e. Develop methods for interpreting public health data that include community input, paying attention to messaging, communication, and narrative. Advance training for the workforce, the public, and communities to use and interpret data.

Context. Representation and power in health decision-making are integral to a modern, equity-oriented public health data system. Power means the authority to shape expectations, decisions, and outcomes in public health. The current public health data system does not prioritize transparency, data access, and use that support sustained civic engagement. Nor does it give deep consideration to the voices of historically marginalized and chronically underrepresented populations, which are required for truth, reconciliation, and racial healing. This focus on equity in data use and decision-making is even more critical in the context of the greater volume, velocity, and variety of data (often used to characterize “big data”), the role of technology in data generation and manipulation (e.g., artificial intelligence, machine learning), and increasing challenges to civic engagement and democracy.

WHAT NEEDS TO BE DONE

SHORT-TERM

- Establish a community advisory council (via the federal advisory committee process) to provide guidance on public health data sources, uses, and interpretation of data and ensure genuine community engagement.
- Work with higher education to develop public health data analytic methods that are interdisciplinary, action oriented, data driven, and aligned with the revised 10 essential public health services.

LONG-TERM

- HHS should provide resources to states, tribes, local health departments, and coalitions to develop best practices for bringing community voice to governance, collection, use case prioritization, and interpretation of data and outcome measures.
- Use a Community Commons (a method of creating a network of changemakers focused on health, equity, and sustainability) model as a data repository to help communities engage with and promote data that are compelling, advance a public health narrative, support action, and demonstrate that health equity can be improved.

WHO NEEDS TO ACT

- HHS
- State and local health departments
- State, regional, county, and municipal planning agencies
- Academia/Research institutions
COMMISSION FORMATION AND DELIBERATIVE PROCESS
Why this Commission Now?

RWJF’s Transforming Public Health Data Systems to Advance Health Equity (TPHD) initiative supports efforts to reimagine the nation’s public health data system to address structural racism and advance health equity. The COVID-19 pandemic has affected communities unevenly across the country, disproportionately burdening people of color in regions, cities, and neighborhoods that have been historically underserved and under-resourced. We know that only by patching together uncoordinated data collection efforts—from local health departments, academic sources, and a host of nontraditional actors—that used different data sources, such as private technology firms, journalist networks, and public intellectuals who have leveraged social media.

For years, a mix of underfunding, complicated governance structures, and technological limitations has plagued these disparate data systems. As a result, their response to COVID-19 was often slow, of varying quality, and unable to answer key questions from public health leaders, policymakers, and the general public. Their failures revealed systemic dysfunction that requires attention, but they also laid bare the impact of structural racism on our capacity to leverage data to improve health. Data have not been consistently collected by race and other demographic characteristics, a reflection of how Black and Brown lives have been undervalued in America. Further, when disaggregated data have been available, they have been interpreted with insufficient contextualization, perpetuating myths and further stigmatizing Black people and communities.

When COVID-19 hit, RWJF launched Transforming Public Health Data Systems to catalyze development of a better public health data infrastructure that would be prepared for the next pandemic and to tackle ongoing public health challenges. While the nation’s current public health data system generally falls short, the COVID-19 pandemic has starkly revealed its failings when it comes to protecting the most vulnerable. For a Culture of Health to truly exist, we need a public health data system that is reliable and effective. And for health equity to advance, we need one that centers populations that historically have been afterthoughts in the conceptualization and design of systems for data collection, sharing, linking, and analysis.

The TPHD Commission was formed in 2020 to review the most significant challenges to the nation’s public health data system and provide recommendations to policymakers, healthcare organizations and institutions, service providers, and philanthropy on how to initiate solutions.

Preparation

Commission deliberations were informed by four complementary activities conducted between October 2020 and April 2021. These components included an environmental scan of literature, reports, coordinated activities, emerging policies, and executive orders related to data modernization and public health data system transformation; and interviews with over 100 individuals selected for their expertise and thought leadership on the main topics to be addressed by the Commission. RAND conducted both efforts between October 2020 and February 2021.

At the same time, RWJF asked a set of research teams (grantees) to conduct targeted studies to deepen understanding of public health data system issues, surface promising practices and case examples of innovation in public health data development and use, and identify systemic changes to facilitate a modern, equity-oriented public health data system. The grantees are continuing their efforts through 2021, but the Commission drew on early insights from their work, as of April 2021. RAND incorporated these insights incorporated into its white papers, which also included findings from the environmental scan and interviews.

The fourth component involved focused discussions with five expert panels on population-specific data gaps (American Indians/Alaska Natives, Blacks/African Americans, LGBTQ+ communities, people
living with disabilities, and women), offering additional insight and context specific to the experiences of these populations. These findings from the expert panels were summarized along with the white papers to inform Commission deliberations.

Each of these components is described in more detail below.

**Four Components of Formative Work**

**ENVIRONMENTAL SCAN**

RAND conducted an environmental scan to identify key issues, points of consideration, tradeoffs and tensions, and current activities related to public health data, data systems, and data modernization efforts. This effort included a targeted scan of published research papers and reports, reviews of websites and working documents describing coordinated activities (e.g., data interoperability), and recent initiatives. It also included a review of federal executive orders, emerging policies, and funding opportunities relevant to public health data systems and health equity, as of June 2021.

The environmental scan initially covered five broad topics: (1) what is public health and public health data, (2) data disaggregation and implications for equity, (3) stewardship and governance of data, (4) private sector innovations in public health data collection and use, and (5) data interoperability. Additional searches included the use of “big data” in public health, data privacy, and ethics of public health data collection. For each topic, the team primarily focused on public health data, but also identified seminal articles and reports from other sectors or disciplines whose findings could apply to public health data systems. For seminal articles, the team reviewed references and pulled additional articles and reports for inclusion in the white papers.

**EXPERT INTERVIEWS**

RAND conducted semi-structured interviews with 112 experts and thought leaders on the main topics before the Commission. Individuals represented diverse sectors, including public health and healthcare, technology and data science, research and policy, journalism, and law. The interviews also included experts in data, data use, equity, community engagement, and research translation who work outside the traditional health sector but in areas germane to the social determinants of health (e.g., environment, community development).

Interviews were designed to elicit insights and perspectives in the following domains, which aligned with the environmental scan. Issues of equity were probed within and across all domains.

- Vision for public health data and gaps and challenges in current public health data, including social determinants of health
- Content and types of public health data
- Data disaggregation and implications for equity
- Access to and use of public health data
- Public health data governance and stewardship
- Role of data providers (public and private sector) and users
- Operational elements of public health data (e.g., features of interoperability, legal issues, privacy)

**RWJF GRANTEES**

Eighteen research teams received grants from RWJF to deepen understanding of public health data system issues and to highlight promising practices and case examples of innovation in public health data systems. Grantee selection was separate from the environmental scan. These in-depth looks are briefly summarized in Table 1 by grant title and theme area. As noted, the grantees are continuing their efforts through 2021. Early insights from each grantee’s work, emerging as of April 2021, were incorporated into white papers developed for the Commission.

The projects may be grouped generally by the themes noted above. Although the grouping below is organized by a primary theme, most grantee work crossed multiple themes and informed the development of the white papers and Commission deliberations broadly.
<table>
<thead>
<tr>
<th><strong>Vision for public health data as well as gaps and challenges in current public health data</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>National Foundation for the Centers for Disease Control and Prevention, Inc.</strong></td>
</tr>
<tr>
<td><strong>Task Force for Global Health, Inc.</strong> (Public Health Informatics Institute)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Content and types of public health data</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>New York University</strong></td>
</tr>
<tr>
<td><strong>United Way Worldwide</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Data disaggregation and implications for equity</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Croal Services Group, LLC</strong></td>
</tr>
<tr>
<td><strong>Urban Institute</strong></td>
</tr>
<tr>
<td><strong>Texas Southern University Foundation</strong></td>
</tr>
<tr>
<td><strong>Drexel University</strong></td>
</tr>
<tr>
<td><strong>University of North Carolina at Chapel Hill</strong></td>
</tr>
<tr>
<td><strong>University of California, Los Angeles</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Access to and use of public health data</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>INFO LINE of San Diego County dba 2-1-1 San Diego</strong></td>
</tr>
<tr>
<td><strong>Health Care Cost Institute, Inc.</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Role of data providers (public and private sector) and users</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>National League of Cities Institute, Inc.</strong></td>
</tr>
<tr>
<td><strong>Community Science, Inc.</strong></td>
</tr>
<tr>
<td><strong>Rush University Medical Center</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Operational elements of public health data (interoperability, legal issues, privacy, ethics)</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Project Evident, a project of the Tides Center</strong></td>
</tr>
<tr>
<td><strong>University of Arkansas for Medical Sciences</strong></td>
</tr>
<tr>
<td><strong>Third Sector New England/Network for Public Health Law</strong></td>
</tr>
</tbody>
</table>
The targeted studies by RWJF grantees surfaced critical issues that the Commission explored further. These topics included the timeliness, quality, and content of data; how data are disaggregated, shared across sectors and organizations, and represented; how community leadership is included in local data decision-making; and how data are translated and used for public health policy.

**POPULATION-SPECIFIC EXPERT PANELS**

The Commission’s formative work included focused discussions with five expert panels on population-specific data gaps (American Indians/Alaska Natives, Blacks/African Americans, LGBTQ+ communities, people living with disabilities, and women). Each panel consisted of individuals who brought expertise to the issues based both on their work and training with data and their own lived experience. Having the expertise and voices from those communities was critical to shaping recommended strategies and approaches to advance data systems and data equity. Each panel convened virtually for eight hours over three days. In addition to panel-specific findings summarized here, all panels highlighted the importance of narrative change around health, well-being, and equity; the need to decolonize and disaggregate data; the importance of authentic community engagement in data decision-making; the importance of training and capacity-building around data; and the need to overlap intersectional identities and experiences in more comprehensive data systems.

**Synthesis of Formative Methods to Guide Commission**

Collectively, findings from these four methodological approaches were synthesized and provided to the Commission in advance of its deliberations. The Commission ultimately had many resources upon which to draw for its work: the series of white papers developed by RAND reflecting the environmental scan, expert interviews, and areas of targeted study; the insights obtained from the expert panels; any additional findings from grantees available at the time of Commission deliberations; and other resources as part of Commission deliberations (described in the next section).

This methodological approach afforded the Commission the depth of multiple perspectives and research streams to inform its recommendations around the need for narrative and structural changes in a modern and transformed public health data system. As noted in the following section, this formative work, combined with deep meeting dialogue and deliberations, distinguished the ultimate Commission recommendations by ensuring that recommended structural changes for the future public health data system are centered on equity, lived experience, and the root drivers of health outcomes.

**Selection of Commission Members and Deliberation Process**

The Transforming Public Health Data Systems Commission process extended over approximately one year and involved several phases leading up to and including the production of the final report. As mentioned, the process began with research and field-scanning work and a portfolio of innovative, action-focused “quick strike” RWJF grants. These Foundation investments provided a set of background reports, level-setting documents, and resources to inform the Commission’s deliberations. These resources also informed the selection process for Commission members.

The process of outreach, vetting, and selection was designed to include diverse scholars, innovators, public health leaders, and activists from across the country. All these phases occurred during the COVID-19 pandemic, which, understandably, constrained the flexibility of some local and national leaders to participate. That said, the resulting panel of Commission members was representative of the
field of public health data research and management, as well as diverse demographics, sectors, and issues.

Given the expressed equity intention of the charge and the urgency created by the pandemic, racial reckoning momentum, and economic upheavals in 2020, the Commission adapted the nationally recognized Truth Racial Healing and Transformation (TRHT) Framework and engagement steps for its deliberations. TRHT is an adaptation to the U.S. context of the globally recognized Truth and Reconciliation (TRC) process, whose most recognized application was in South Africa. Envisioned by Gail C. Christopher and launched in 2016 by the W. K. Kellogg Foundation in collaboration with several other philanthropies, TRHT is now used by thousands in colleges and universities, nonprofits, government, and private entities around the United States. This framework has five pillars:

- Narrative Change
- Racial Healing and Relationship Building
- Separation
- Law
- Economy

FIGURE 1. TRHT FRAMEWORK
These pillars reflect the comprehensive, coordinated work and strategies required for collective movement toward a more racially just and equitable society. Our stories about who we are as a country and how we became America must be revisited and revised to reflect more authentic, expansive, and inclusive voices of diverse groups. At the same time, we must build individual and collective capacities and tools for relating across perceived barriers and entrenched lines of difference. This requires relational skills and competencies associated with racial healing and cross-identity group communication.

These processes of changing racial hierarchy narratives and expanding circles of engagement for equity and justice enable the work on the remaining pillars of the TRHT Framework: Separation, Law, and Economy. These three words embody the primary systemic and structural tools and means used to create and sustain systems of exclusion, oppression, and racial hierarchy based on the belief in a hierarchy of human value. Now that racism has been declared a “public health crisis” by more than 200 local health jurisdictions and, most recently, by the CDC, it is time to transform public health data systems into engines that can help drive the healing and transformation of our democracy from its centuries-old roots in racial hierarchy toward its aspirational tenets of equal human rights and fairness for all.

The TRHT Framework requires five sequential steps. The first is visioning and reimagining the successfully changed future. This is done for each house pillar. Once the future state is agreed upon, participants analyze the current landscape or “now” in light of this vision. This step is followed by identifying resources, human and financial, needed to achieve the vision. Finally, participants develop short-term and long-term recommendations for specific actions by designated institutions, sectors, or groups. Members of the TPHD Commission, together and in teams representing the five pillars, undertook this process.

Charged with reimagining the nation’s public health data system to address structural racism and advance health equity, the Commission was convened virtually and engaged in a participatory deliberation process designed and facilitated by the National Collaborative for Health Equity (NCHE) team. Teams aligned with the five TRHT pillars discussed their visions, deliberated priorities, and shaped recommendations collaboratively. Ultimately, the visioning and recommendations from the five TRHT Framework pillars were adapted and condensed into three focus areas for public health data and data systems transformation efforts, reflecting the vision statements developed by the Commission teams as they deliberated.

The first theme, **Health Equity and Well-Being in Narrative Change**, was reflected in vision statements calling for sustainable and transparent data collection conducted in collaboration with communities and people most impacted by social and economic disparities to build trust and provide the contextual and historical data needed to change the narrative, disrupt the status quo, and enable restorative and prospective storytelling.

The second theme, **Equitable Governance, Systems, and Community Engagement**, emerges from vision statements related to building an agile, well-resourced public health data system that respects communities’ agency in understanding and governing themselves and has a high degree of interoperability, allowing for stakeholder engagement and interaction. The transformed public health data system would take a broad view of both the processes and outcomes of health and well-being, encompass different domains of life (e.g., health, economy, social, cultural, environmental) across systems and institutions (e.g., education, housing, labor, development, health care, criminal justice, immigration); measure both harmful exposures and positive assets/strengths/opportunities of communities, places, and experiences; and account for upstream macrostructural factors (policies, practices, and systems).

The **Measuring and Addressing the Health Impact of Structural Racism and Other Inequities** theme reflects vision statements about the need for a public health data system that characterizes separation, segregation, and barriers to opportunities in
measurable ways, and addresses the problems of missing data and misuse of data. The data system would set new targets and measure progress toward those targets, addressing needs for storytelling, data analytics, policy development and evaluation, and programmatic use. It would monitor wellness for the whole person and the whole community and help determine what works and what does not.

Conclusion

Time and again, major crises such as global pandemics precipitate meaningful and constructive societal change. Sewer systems, public parks, and clean water systems are a few examples of major innovations that transformed cities and society following pandemics. It seems fitting that, in this information era, the COVID-19 pandemic will generate innovation and transformation in how public health data are collected and leveraged to foster better health and well-being for all.

We hope that the recommendations in this report will help transform understanding of public health data and how they can be applied to advance health equity and racial justice. COVID-19 revealed stark and painful racial inequities in the determinants of health and opportunities for well-being. In so doing, this pandemic and its attendant racial reckoning and economic crises must compel actions by and across all sectors to address and redress historic and contemporary patterns of inequitable access to opportunity. COVID-19 demonstrated the shared costs of inequality. We now have greater insight into the role of accurate, timely, accessible, and comprehensive public health data in driving decisions for needed investments to produce better and more equitable outcomes—whether it’s access to safe, affordable, stable housing or transportation to and from vaccination centers.

Public health data can be used to create unprecedented systems of accountability for progress toward greater fairness and equity. The recommendations in this report aim to engage and catalyze representatives from local communities, nonprofit organizations, and academia, as well as from private, public, and philanthropic sectors. It is a report for and call to action to the nation about how we must collaborate, expand, and innovate to measure and invest in the health and well-being of all populations.
COVID-19 and the national reckoning with racism and other social inequities have activated a long overdue national health consciousness. Critical to the success of the modern public health data system is a focus on equity in the story, content, and structure of public health data; how those data are used to address the painful toll of racism and other systemic inequities; and the engagement of diverse stakeholders in access to and use of data for decision-making—all with attention to issues of power, marginalization, and justice. Now is the time for the public health field to crystallize a “moonshot” or a more ambitious north star to chart a clear path to health, well-being, and equity that enables everyone to live their healthiest life and thrive. This shared vision can help sectors and stakeholders accelerate action, commit to transforming the public health data system through their own work, and join forces to amplify impact. This section outlines the ongoing actions that should be pursued by government as well as private and other civil society sectors to transform our public health data system.

Government

Federal, state, and local government leaders can support critical public health data infrastructure, clarify the need for consistency in public health data (e.g., how subgroups are captured), and affirm how data can help address structural racism and other inequities. Here are examples by level of government:

FEDERAL GOVERNMENT

- Develop minimum standards for data collection, disaggregation, presentation, and access in federally funded data collection efforts, with an orientation to “freeing federal data” or promoting greater access. Standards should include how to balance privacy considerations, the importance of subgroup analyses, the critical application of relevant tribal policies in data review, how the impact of racism is measured, and federal workforce training in public health data.
- Strengthen public health data infrastructure and incentivize the use of new data collection and analytic approaches through a public messaging campaign and a combination of policies, funding mechanisms and priorities, and guidance to state and local agencies.
- Restrict federal funding for data infrastructure to systems that are standards-based and interoperable.
- Offer guidance on interpretation of racial/ethnic variation in health-relevant data to counter longstanding acceptance of the idea that these variations reflect biological differences rather than systemic, cultural, behavioral, and social factors.

STATE GOVERNMENT

- Ensure that state policies for public health data collection, sharing (including balancing privacy and transparency), and analysis are equity driven and explicitly call out the influence and consequences of structural racism and other inequities on health.
- Partner with local health departments and departments that provide public health data (e.g., social services data) to consider new models of collaboration to improve efficiency and timeliness of decision-making and action at state and local levels.

LOCAL GOVERNMENT

- Regularly connect public health data to local communications (i.e., what is happening in the community and how it relates to overall community well-being, or the impact of inequity).
- Ensure that local voice is represented not only in public health data (e.g., from whom data are collected), but in those positions of authority responsible for making sense of the data and informing decisions.
- Explore data-sharing collaborations across government and civil society (e.g., nonprofits, businesses) that can more consistently generate public health data to support equity considerations and advance innovative public-private collaborations on data and analysis.
Public Health

Eroding public trust and sustained disinvestments in public health have produced significant variability in capacity and infrastructure to collect, analyze, share, and leverage existing data to identify and address emerging health concerns and structural inequities that affect health.

- Lead multisector collaboration around public health data sharing to improve the timeliness and quality of data to strengthen local decision-making.
- Strengthen capacity, diversity, and ongoing training of the public health workforce to monitor and address health equity, both in the field of public health and through novel collaborations with business, academia, or other sectors that influence health.
- Advocate for and prioritize modernization efforts and data sharing within and across the public health system to ensure that local data can inform emerging public health concerns at the regional, state, and national levels in real time.

Healthcare Systems

Healthcare systems have increased the collection of data on SDOH and other “non-clinical” factors that influence health (e.g., isolation, lack of social support), primarily at the individual level.

- Consistently collect SDOH data during consumer encounters, using standardized questions and ICD-10 codes that allow data aggregation within communities.
- Connect these data on individual social needs (e.g., housing), with structural inequities confronted by consumers (e.g., neighborhood characteristics, trauma exposure, experiences of discriminatory policies) to offer useable information on equity to the community.
- Overcome historic silos and build partnerships and legal solutions to facilitate sharing of relevant healthcare data with public health departments in a timely and efficient manner, allowing data linkages and disaggregation of subgroups and geographic regions.

Businesses

Businesses generate and analyze health data. Very few of these data, however, are consistently used for the public good to identify emerging health needs or to inform local decision-making.

- Work with government partners and other organizations to develop standards through which public health data generated by the private sector can be used and communicated.
- Engage in more consistent workforce exchanges and data partnerships with public health organizations to cross-pollinate innovations in the types, content, quality, and precision of public health data.
- Foster public health innovation by promoting methods in use by technology companies, such as data integration, “big data” analytics, and data security.

Nonprofits, Professional Associations, and Schools

Nonprofits and community-based organizations are foundational to understanding existing and emerging health needs of the community. Professional associations, including the Association of State and Territorial Health Officials and the National Association of County and City Health Officials, and research institutions like the National Academies of Sciences, Engineering and Medicine are trusted organizations and often the go-to places for information and resources for their members. Schools of public health are developing the future workforce for the modern public health data system.
NONPROFITS

- Build trust in data as a public good in the community and among constituents, identifying relevant issues (e.g., around privacy or how data will be used). Support the development of solutions that address concerns to ensure that such data do not further marginalize or bring harm to populations.
- Work locally to develop data-sharing strategies with government, and, where appropriate and where there is health benefit, reciprocity, and equitable access.

PROFESSIONAL ASSOCIATIONS

- Develop strategies to improve data completeness and quality, particularly with respect to equity and drivers of health and well-being.
- Actively work with professional associations from other sectors that influence health (e.g., education, social services, environment) to align efforts, particularly around standardization and data use. This can minimize variability across sectors that impedes effective data sharing and understanding around health trends at a local level.

SCHOOLS

- Help students monitor and evaluate structural and systemic inequities and understand what data systems must include to call out those inequities.
- Train the next generation of public health practitioners and researchers to meaningfully partner with local stakeholders and to value lived experience and community expertise as much as formal training.

PHILANTHROPY

Philanthropic organizations can guide an agenda by calling out important data gaps (e.g., positive health and well-being, structural inequity data), and offer strategic and catalytic investments to support methodologic advancements and generate new solutions.

- Promote a national, unifying health goal that can be understood and executed, with a focus on the role of structural racism and the values of health equity and well-being.
- Leverage convening and funding opportunities to meaningfully engage leaders, including those from non-health sectors and community organizations, who can advance the importance of public health data.
- Focus funding on key parts of data systems modernization, such as improving governance and using new forms of data and analytic methods to convey better stories of health equity and well-being.
ACKNOWLEDGEMENTS

The preparation of this report, like the orchestration of the TPHD Commission, was a collective effort. The work of staff and consultants from multiple organizations is deeply appreciated. These groups include National Collaborative for Health Equity, Virginia Commonwealth University Center on Society and Health, RAND Corporation, MMS Education, and Burness. These collective efforts were made possible by the vision and generous support of the Robert Wood Johnson Foundation.

GLOSSARY

CONFIDENTIALITY. The obligations of individuals or groups that receive or use information to respect the privacy interests of individuals who are subjects of the data.

DATA DISAGGREGATION. The collection, provision, and reporting of data by subcategories.

DATA EQUITY. The collection, provision, and reporting of demographic data that consider the equity consequences and sociopolitical nature of the category definitions.

EQUITY. Defined in shorthand as the fair and just access to opportunity. There are many types of equity, such as:

- Distributive equity focuses on allocation and resource management decisions, with attention to the balance of costs, risks, and benefits. Usually, distributive equity considers how decisions are made and benefits are distributed based on dimensions of need and social benefit.

- Procedural equity addresses how the concept of fairness is included in approaches and policies once the equity parameters are set (i.e., for whom are we improving equity).

- Contextual equity is the backdrop for both procedural and distributive equity, accounting for the political, economic, social, and intergenerational factors that influence how populations engage with society and its systems and benefits. This includes contextual variables such as access (e.g., access to capital) and power (e.g., the ability to gain and maintain access to resources).

EQUITY-ORIENTED. A public health data system that centers equity, or the fair and just access to the opportunity to be healthy, in the content and structure of data; the engagement of diverse stakeholders in the access to and use of data for decision-making, with attention to issues of power, marginalization, and justice, and in how data are used to address systemic inequities.

HEALTH EQUITY. Everyone has a fair and just opportunity to be as healthy as possible. This requires removing obstacles to health such as poverty, discrimination and its consequences, powerlessness, as well as lack of access to good jobs with fair pay, quality education and housing, safe environments, and health care (Braveman, 2017).

HEALTH INEQUITIES. Systematic differences in the opportunities that groups have to achieve optimal health, leading to unfair and avoidable differences in health outcomes. The dimensions of social identity and location that organize or “structure” differential access to opportunities for health include race and ethnicity, gender, employment and socioeconomic status, disability and immigration status, geography, and more.

INTEROPERABILITY. The ability of systems that create, exchange, and consume data to have clear, shared expectations for the contents, context, and meaning of those data, ensuring that information is shared appropriately among systems and partners in the right format, through the right channel, and at the right time (www.cdc.gov and Data Interoperability Standards Commission).
GOVERNANCE. The structures and processes by which people in societies make decisions and share power, creating the conditions for ordered rules and collective action. Data governance refers to how data are structured, shared, and protected.

MACHINE LEARNING. The use of computer systems to learn and adapt by using algorithms and other models to analyze and draw inferences from data patterns.

NARRATIVE. Narratives shape the way people see and think about the world around them. They are expressed, received, and internalized through stories and experiences in art, popular culture, traditions and common practices, the built environment, policies, systems, and structures, and more. Aggregated over time—and filtered through lived experience, culture and environment, and the echo chambers where people seek feedback and validation—narratives influence the way people make sense of their surroundings, interpret information, and make decisions. (See metgroup.com/narratives for more information.)

PARTICIPATION. The active role of those impacted by public health decisions in the process of arraying decision options and providing meaningful input on those decisions.

POWER. The authority to shape expectations, decisions, and outcomes in public health.

PRIVACY. An individual’s rights to control the acquisition, uses, or disclosures of his/her/their identifiable health and other data.

PUBLIC HEALTH DATA SYSTEM. The Commission defines a modern, transformed public health data system as one that is accountable to and reflects the perspectives of local communities and diverse populations, including people of color, immigrants, persons with disabilities, and the LGBTQ+ community. The system must be sustainable, fully integrated with healthcare and other sectors that drive health by linking data across multiple sectors and at multiple levels, and provide data that are disaggregated, comprehensive, and timely. It should move from problem-focused to solution-focused research and have the capacity to provide the knowledge needed to optimize health and well-being for all people. It must provide tools to address racism and racial/ethnic and other social disparities; it should facilitate restorative systems that respect all communities, including the sovereign right of tribal nations to govern data and ensure they have access to them; protect individuals’ privacy and security; and guard against unintended consequences.

SECURITY. The technological or administrative safeguards or tools designed to protect identifiable health data from unwarranted access or disclosure.

SENSEMAKING. The cognitive processes by which people make meaning from data and experiences.

SMALL AREA ESTIMATES. Using statistical techniques to develop estimations of small sub-populations.

STRUCTURAL RACISM. The laws, policies, cultural representations, and norms across interconnected systems that support the unfair treatment of some groups based on the social construct of race that is grounded in a false ideology of a hierarchy of human value.

SYNTHETIC DATA. Information that is artificially generated rather than produced from actual events.

VOICE. The inclusion of the perspectives, ideas, and lived experiences of those impacted by public health decisions.
REFERENCES


20. Technology TOotNCfHI. 2020-2025 Federal Health IT Strategic Plan. Office of the Secretary, United States Department of Health and Human Services; 2020.

21. Technology TOotNCfHI. ONC’s Cures Act Final Rule supports seamless and secure access, exchange, and use of electronic health information. HealthIT.


RESOURCES

Business Case for Racial Equity

Closing the Racial Inequality Gaps: The Economic Cost of Black Inequality in the U.S.
https://irciti.com/%2FPRxPvgyNWu3l9AU1aiGr%2BsK-bjlijBjSaTOSdw2DF4xynPwFB8a2jV1FaA3Idy7vY59b0t-N2lxVQM%3D

Commission on Security and Cooperation in Europe (Helsinki Commission), Briefing on Truth, Reconciliation, & Healing: Toward a Unified Future

Deconstructing Inequities — Transparent Values in Measurement and Analytic Choices

Ethics and Empathy in Using Imputation to Disaggregate Data for Racial Equity: Recommendations and Standards Guide

The HOPE Initiative: Measures to Advance Health and Opportunity
https://www.hopeinitiative.org/

Including disability in all health equity efforts: an urgent call to action
https://www.thelancet.com/journals/lanpub/article/PIIS2468-2667(21)00115-8/fulltext

Physician–patient racial concordance and disparities in birthing mortality for newborns

Racism and Health: Evidence and Needed Research

The Impact of Chronic Underfunding on America’s Public Health System: Trends, Risks, and Recommendations

Truth, Racial Healing, and Transformation: Creating Public Sentiment

Understanding Hate Through a Public Health Lens
MEET THE COMMISSIONERS

GAIL C. CHRISTOPHER, DN, is the director of the National Commission to Transform Public Health Data Systems and serves as executive director of the National Collaborative for Health Equity. She is also a senior scholar at George Mason University’s Center for the Advancement of Well-Being. Christopher is an award-winning social change agent with expertise in the social determinants of health and well-being and in related public policies. She is known for her pioneering work to infuse holistic health and diversity concepts into public sector programs and policy discourse. In her role as the senior advisor and vice president at the W.K. Kellogg Foundation (WKKF), she was the driving force behind the America Healing initiative and the Truth, Racial Healing and Transformation effort. Christopher also served as WKKF’s vice president for programs. In 2015, she received the Terrance Keenan Award from Grantmakers in Health. She chairs the Board of the Trust for America’s Health. She is the visionary for and architect of the WKKF-led Truth Racial Healing and Transformation (TRHT) effort for America. TRHT is an adaptation of the globally recognized Truth and Reconciliation Commission (TRC) model.

MARGARITA ALEGRIÁ, PHD is chief of the Disparities Research Unit at Massachusetts General Hospital, and a professor in the Department of Psychiatry at Harvard Medical School. Alegría is currently the principal investigator of four research studies funded by the National Institutes of Health: The Impact of Medicaid Plans on Access to and Quality of Substance Use Disorder Treatment, Building Infrastructure for Community Capacity in Accelerating Integrated Care, Building Community Capacity for Disability Prevention for Minority Elders and Latino Youths in Coping with Discrimination: A Multi-Level Investigation in Micro- and Macro- Time.

MARY T. BASSETT, MD, MPH, was recently appointed as the Health Commissioner for the state of New York. Bassett has dedicated her career to advancing health equity. Dr. Bassett currently serves as the director of the François-Xavier Bagnoud (FXB) Center for Health and Human Rights at Harvard University and the FXB professor of the Practice of Health and Human Rights at the Harvard T.H. Chan School of Public Health. Prior to joining the FXB Center, she served as New York City’s commissioner of health from 2014 to 2018.

RAYMOND BAXTER, PHD, is co-chair of the National Academies of Science, Engineering, and Medicine (NASEM) Roundtable on Population Health, serves on the CDC Foundation Board of Directors, is a trustee of the Blue Shield of CA Foundation, and serves as an advisor to the deans of the University of California Berkeley School of Public Health and the University of California San Francisco School of Nursing. He most recently served as president and CEO of Blue Shield of California Foundation, leading its mission to make California the healthiest state and end domestic violence by addressing the root causes of ill health and inequity. For 15 years, Baxter was Kaiser Permanente’s senior vice president for community benefit, research, and health policy. Previously, he headed the San Francisco Department of Public Health, the New York City Health and Hospitals Corporation, and The Lewin Group.

JULIET K CHOI, JD, is chief executive officer of the Asian and Pacific Islander American Health Forum, a national health justice organization that influences policy, mobilizes communities, and strengthens programs and organizations to improve the health of Asian Americans, Native Hawaiians, and Pacific Islanders. She is an accomplished cross-sector leader and coalition builder who specializes in change management, system reform, and stakeholder relations, particularly in the areas of immigration, civil rights, healthcare, and disaster relief.
MICHAEL CRAWFORD, MBA, MHL, is the associate dean for strategy, outreach, and innovation at the Howard University College of Medicine and founding executive director of Howard University’s 1867 Health Innovations Project. Prior to Howard University, Crawford served as the chief of staff at Unity Health Care, Inc., one of the largest health center networks in the United States. Prior to Unity, Crawford held domestic and international leadership positions at Johnson & Johnson, GlaxoSmithKline, and Gannett Company. Crawford brings expertise at the intersection of digital health equity, data, strategy, product development, policy, and operations. Crawford’s work focuses on developing scalable digital health and data models to help enhance health access, outcomes, and affordability for medically underserved and vulnerable populations.

FERNANDO DE MAIO, PHD, is the director of research and data use for the Center for Health Equity at the American Medical Association and a professor of sociology at DePaul University. His research and teaching interests lie primarily within medical sociology and social epidemiology, with a focus on the concept of structural violence. His work has been guided by the notion of “radical statistics”—the idea that statistical analysis can be used to not just describe the world, but to change it. He is the author of Global Health Inequities (Palgrave Macmillan, 2014) and co-editor of Community Health Equity: A Chicago Reader (University of Chicago Press, 2019) and Unequal Cities: Structural Racism and the Death Gap in America’s 30 Largest Cities (Johns Hopkins University Press, 2021).

KAREN DESALVO MD, MPH, MSC, is the chief health officer at Google. DeSalvo served as acting assistant secretary for health at the U.S. Department of Health and Human Services in the Obama administration. Under her leadership, HHS set and met historic goals in payment reform, supported transformed models of care delivery, including in primary care, and changed the approach to information distribution in the health system. She also served as the National Coordinator for Health Information Technology, where she set national strategy and policy on health information technology and championed interoperability in health settings.

ABIGAIL ECHO-HAWK, MA (PAWNEE), is the executive vice president of the Seattle Indian Health Board and the director of the Urban Indian Health Institute, a tribal epidemiology center. She works to support the health and well-being of urban Indian communities and tribal nations across the United States. Echo-Hawk has been recognized as a national leader in decolonizing data for Indigenous people, by Indigenous people.

THOMAS LA VEIST, PHD, is dean of the School of Public Health and Tropical Medicine at Tulane University in New Orleans, LA. Before joining Tulane, LaVeist was chairman of the Department of Health Policy and Management at the George Washington University, Milken Institute School of Public Health, and spent 25 years on the faculty of the Johns Hopkins Bloomberg School of Public Health. LaVeist’s research focuses on the development of policy and interventions to address race disparities in health-related outcomes.

ALEXIS C. MADRIGAL is a writer at The Atlantic and the co-founder of the COVID Tracking Project. He’s been a visiting scholar at the University of California Berkeley’s Information School as well as the Center for Science, Technology, Medicine, and Society.
JOHN LUMPKIN MD, MPH, is president of the Blue Cross and Blue Shield of North Carolina Foundation, since April 2019. He leads the organization in pursuit of its stated mission to improve the health and well-being of everyone in North Carolina by focusing on transforming the health care system (including oral health), expanding access to healthy food, supporting a healthy start in life for children, improving the physical conditions where people live, and strengthening the ability of communities to improve health.

AMY O’HARA PHD, MA, is a research professor in the Massive Data Institute and executive director of the Federal Statistical Research Data Center at Georgetown University’s McCourt School for Public Policy. She also leads the Administrative Data Research Initiative, improving secure, responsible data access for research and evaluation, and is co-founder of the Civil Justice Data Commons. O’Hara addresses risks involved with data sharing by connecting practices across the social, health, computer, and data sciences. Her research focuses on population measurement, data quality, and record linkage. Prior to joining Georgetown, O’Hara was a senior executive at the U.S. Census Bureau, where she founded the administrative data curation and research unit.

JONATHAN PERLIN MD, PHD, is president, clinical operations and chief medical officer at HCA Healthcare, where he leads a team in using a learning health system model for improving care at the system’s 185 hospitals and 2,200 sites of care. The effort achieved national recognition for preventing elective pre-term deliveries, reducing maternal mortality, increasing sepsis survival, and developing public-private-academic partnerships for improving infection prevention and treating COVID-19. Prior to HCA, Perlin was under secretary for health in the U.S. Department of Veterans Affairs. He is a MedPAC commissioner, a Congressional Budget Office health advisor, chairs the National Quality Forum, and is an elected member of the National Academy of Medicine. He has faculty appointments at Vanderbilt University and at Virginia Commonwealth University.

NINEZ PONCE MPP, PHD, is a professor in the University of California Los Angeles (UCLA) Fielding School of Public Health, director of the UCLA Center for Health Policy Research, and principal investigator for the California Health Interview Survey. Her research contributes to the elimination of racial/ethnic/social disparities in health. Ponce recently served on the Board of Scientific Counselors, National Center for Health Statistics. She has served on committees for the National Academy of Sciences and the National Quality Forum, where her expertise has focused on setting guidance for health systems in the measurement and use of social determinants of health as tools to monitor health equity. In 2019, Ponce and her team received the AcademyHealth Impact award for their contributions to population health measurement to inform public policies.

CHESLEY RICHARDS, MD, MPH, FACP, served at the Centers for Disease Control and Prevention (CDC) from 1998 to 2020 in several roles, including as deputy director for public health science and surveillance. In this position, he was responsible for strengthening CDC’s science foundation by working across the Office of Science, the Office of Laboratory Science and Safety, the Center for Surveillance, Epidemiology, and Laboratory Services, and the National Center for Health Statistics. A primary focus of his role was to advance an agency-wide public health data strategy and serve as an advisor to the CDC director.

JA VIER ROBLES, JD, is a faculty member and professor of the Kinesiology and Health Department and is the director of the Center for Disability Sports, Health and Wellness at Rutgers University. He is the chair of the New Jersey Disabilities Covid-19 Action Committee and was appointed by Gov. Murphy to the Puerto Rico commission. Robles is a board member of the United Spinal National Board and the vice president of the Latino Action Network of New Jersey. He is past president of Thisabled, LLC, an organization that provides support to persons with disabilities through self-empowerment and perseverance. He is the founder of the Facebook group “People with Disabilities Helping Each Other Survive the Coronavirus.” Robles has written for numerous publications including, Latinos NJ, ThisAbled Nation, and New Mobility. One of his poems was recently published in the book Access Granted.
The National Commission to Transform Public Health Data Systems was convened by The Robert Wood Johnson Foundation.