What Gets Measured Gets Done: An Assessment of Local Data Uses and Needs in Large Urban Health Departments

Brian C. Castrucci, MA; Elizabeth K. Rhoades, PhD; Jonathon P. Leider, PhD; Shelley Hearne, DrPH

Context: The epidemiologic shift in the leading causes of mortality from infectious disease to chronic disease has created significant challenges for public health surveillance at the local level. Objective: We describe how the largest US city health departments identify and use data to inform their work and we identify the data and information that local public health leaders have specified as being necessary to help better address specific problems in their communities. Design: We used a mixed-methods design that included key informant interviews, as well as a smaller embedded survey to quantify organizational characteristics related to data capacity. Interview data were independently coded and analyzed for major themes around data needs, barriers, and achievements. Participants: Forty-five public health leaders from each of 3 specific positions—local health official, chief of policy, and chief science or medical officer—in 16 large urban health departments. Results: Public health leaders in large urban local health departments reported that timely data and data on chronic disease that are available at smaller geographical units are difficult to obtain without additional resources. Despite departments’ successes in creating ad hoc sources of local data to effect policy change, all participants described the need for more timely data that could be geocoded at a neighborhood or census tract level to more effectively target their resources. Electronic health records, claims data, and hospital discharge data were identified as sources of data that could be used to augment the data currently available to local public health leaders. Conclusions: Monitoring the status of community health indicators and using the information to identify priority issues are core functions of all public health departments. Public health professionals must have access to timely “hyperlocal” data to detect trends, allocate resources to areas of greatest priority, and measure the effectiveness of interventions. Although innovations in the largest local health departments in large urban areas have established some methods to obtain local data on chronic disease, leaders recognize that there is an urgent need for more timely and more geographically specific data at the neighborhood or census tract level to efficiently and effectively address the most pressing problems in public health.

KEY WORDS: BCHC, informatics, informatics needs, information, public health data

Industrialization and immigration resulted in overcrowded housing and a general lack of sanitation in large urban areas throughout the United States in the 1900s. These conditions contributed to repeated outbreaks of infectious disease. Local health departments (LHDS) in these large urban areas responded to this challenge with innovations and improvements to sanitation, vaccination programs, and disease surveillance; this significantly reduced infectious disease morbidity and mortality. In the early 1900s, for example, New York City had at least 8 major epidemics of cholera, dysentery, smallpox, and diphtheria, which together took more than 25,000 lives. However, by 1925,
the city had virtually eliminated deaths due to these communicable diseases. These accomplishments were possible, in part, because local health officials were able to collect and use data about specific practices in specific places. During the 20th century, advances in laboratory techniques and technologies, improved screening tests, and more accurate disease reporting translated into faster and better responses to disease outbreaks.

While control of communicable diseases—both known and emerging—continues to be a priority, the increased prevalence of chronic disease presents new challenges for LHDs. Chronic disease accounted for 4 times the proportion of all US deaths in 2000 compared with those in 1900. By 1925 in New York City, mortality rates for cancer and heart disease were more than 175%, which is greater than that just 50 years earlier. By 1925, the mortality rate associated with diseases of the arteries had already increased 650% compared with that of the late 1800s. The top 3 causes of death in 1900—pneumonia, tuberculosis, and diarrhea and enteritis—have been replaced by heart disease, cancer, and stroke.

As disease dynamics have changed and the role of personal behaviors in premature mortality has been more widely recognized, surveys—such as the Behavioral Risk Factor Surveillance Survey (BRFSS; piloted in 1981, initiated nationally in 1993), Youth Risk Behavior Survey (initiated in 1991), Pregnancy Risk Assessment and Monitoring System (initiated in 1988), National Health and Nutrition Examination Survey (first conducted in 1971, done annually since 1999)—were implemented to provide state-level data to aid agencies trying to target resources to reduce behavioral risks and their consequent illnesses. The emergence of these state-level surveys demonstrated an understanding that national data may not be applicable to the conditions found in any given state. A similar argument could justify the need for data at the substate and subcounty levels—state- or county-level data may not be applicable to the conditions in a given community or neighborhood. However, for methodologic reasons, these surveys rarely provide any data below county levels, such as data from neighborhoods, census tracts, block groups, or other common subcity geographic jurisdictions. States and even some cities have implemented corresponding efforts to collect estimates of disease prevalence and correlates below the county level, but few mechanisms were developed that had the geographic precision and clinical accuracy necessary to monitor chronic conditions at the local level.

The relatively recent expansion of health information technology (IT) has presented new opportunities for the systematic collection and analysis of large-scale health care data to inform public health practice. In 2008, 9.4% of all nonfederal acute care hospitals had adopted at least a basic electronic health record (EHR) system. Federally funded incentives have contributed to a 5-fold increase in at least basic EHR adoption from 2008 to 2013. The increase in EHR adoption along with the proliferation of other electronically captured, digitized health information has increased the availability of health care data. However, it is unclear how public health leaders will capitalize on this new source of information.

Today, leaders in large urban health departments are developing strategies to meet the critical need for precise and timely local data to inform decision-making both to improve public health and to use evidence-based policies to inform their work. These strategies have not been documented systematically, nor have the opinions of this group in relation to ongoing data needs that are required to ensure the continued future impact of LHDs. To gather this information, we interviewed local health officials and senior staff members—including scientific leads—who are responsible for the health of some of the country’s largest municipalities, about their perspectives on data, their achievements to date in getting and using data to inform their public health strategies, and their views on the challenges and concerns they now face.

**Methods**

This analysis was part of a larger project that gathered perspectives on critical public health issues from leaders of the Big City Health Coalition (BCHC), a group representing 20 of the largest health departments. The BCHC LHDs cover 46 million people, or 15% of the US population. Detailed methods of the larger study have been described elsewhere. For this analysis, we used a mixed-methods design that included interviews with key informants and a smaller survey to capture organizational characteristics related to data capacity.

Between August and October 2013, we conducted semistructured interviews using a single interviewer with 45 leaders from 16 participating LHDs in the BCHC. The instruments were pretested with 5 current and former public health practitioners. These participants held 1 of 3 positions in each LHD—local health official, chief of policy, or chief science/medical officer. All 3 positions were not represented by every BCHC, as each agency does not necessarily have a chief science/medical officer or a chief of policy. Interview questions were open-ended and focused primarily on data needs, capacities, perceived achievements, and predictions about the future of public health informatics. Qualitative data were independently coded by 2 of...
the authors and were managed and analyzed in NVivo 10 (QSR International, Cambridge, Massachusetts). A preliminary codebook was developed on the basis of major sections of the interview instrument. The codebook was used independently by 2 researchers to code the interviews in 2 groups. After each round of coding, researchers compared results, disagreements in coding were resolved, and data were recoded using a finalized codebook. Data were analyzed in aggregate, as well as by department location.

All 45 participants took part in the open-ended interviews. However, only the chief science/medical officers were asked to complete the Web-based survey portion of the project, which asked about data and informatics capacities in their departments, since staff in these positions have the most experience and expertise in this aspect of health department work. We integrated the data during analysis, per the embedded mixed-methods design. This involved linking survey responses to interviewee and conducting several qualitative analyses with, for example, organizational capacities or levels of data use/sophistication in mind. We report these integrated results later. We pretested all instruments used in this study with 5 current and former public health practitioners. We report descriptive statistics below. Quantitative data were cleaned, managed, and analyzed in Stata 13 (StataCorp LP, College Station, Texas).

Results

LHD health IT and informatics capacity: Survey results

Survey questions given to health departments’ chief science/medical officers (n = 12/14 responded) asked about how IT was used in their departments to collect and disseminate data. These departments more often collected data related to traditional public health concerns, such as monitoring of infectious disease and food inspections; collecting data on varying contemporary health indicators, such as cancer or chronic disease was much less common.

Less than half of the LHDs used systems that collected or disseminated data from EHRs or health information exchanges (HIEs). Participants also indicated that there was considerable variation between departments in systems capability for bidirectional reporting and exchange, as well as whether systems were able to access and use local or state data.

Prevalence of electronic data collection

Figure 1 shows the use of electronic data collections by the study participants. Twelve departments most commonly used electronic systems to collect data about control and prevention of communicable diseases. Eleven of 12 responding BCHC LHDs used electronic disease reporting systems, 10 used immunization registries, and 9 used electronic laboratory reporting. Few departments used electronic systems for disease registries: 2 had electronic cancer registries, and 7 had electronic registries for other diseases. At the program level, use of electronic data was more common in traditional public health areas such as (eg, reportable diseases, laboratory results, and case management). Few departments had data on food service inspections, mapping, and EHRs, or on management of disease outbreaks.

Use of HIEs

The use of HIEs was not widespread (Figure 2). Half of the LHDs reported that they had no interaction with HIEs, whereas the other half used HIEs for a variety of data, typically related to collecting and analyzing/integrating data on emerging infectious diseases. The HIEs were less commonly used for monitoring indicators of health care quality, chronic disease, or health disparities. Two departments reported using HIEs to monitor risk factors for chronic disease, such as smoking, physical activity, and diet; only one reported using an HIE for syndromic surveillance and another for environmental exposures. Only a few LHDs reported using HIEs to collect data, with several reporting use of HIEs for laboratory results and reportable diseases. Fewer than 3 respondents reported using HIEs for any other program activity, including EHRs, hospital discharge data, chronic disease monitoring, vital records, and the Supplemental Nutrition Program for Women, Infants, and Children.

Capability for bidirectional reporting and exchange

Not all electronic systems are capable of sharing data bidirectionally, where each system is able to send and receive information from the other. Overall, this capability is uncommon among responding BCHC LHDs. Bidirectional exchange in health-related electronic data systems generally has developed following the pattern of electronic data collection overall, with 2-way exchange of data about reportable diseases and laboratory results coming first. While 12 departments reported that they collected information electronically on reportable diseases, only 9 had the capability for 2-way data exchange. Five of 10 departments that collected electronic data on case management could share information bidirectionally. Both of the 2 cancer registries were capable of bidirectional data exchange, but only 4 of 11 departments had the same capability with electronic laboratory reports on communicable diseases (Figure 3).
FIGURE 1  Programs’ Use of Electronic Data Collection (N = 12)

- Reportable diseases
- Laboratory results
- Case management
- Food service inspections
- Outbreak management
- Geographic coded data for mapping analysis
- Electronic health record (personal health services)
- Vital records
- Medicaid billing
- Maternal and child health reporting
- Chronic diseases
- Other health care systems data (eg, bedavailability)
- Hospital discharge data
- WIC
- On-site waste water treatment systems
- Water wells (licensing and/or testing)

Abbreviations: HIE, health information exchange; WIC, Supplemental Nutrition Program for Women, Infants, and Children.

FIGURE 2  Cities’ Use of Health Information Exchange (N = 12)

- None
- Emerging infectious diseases
- Health care quality indicators
- Chronic disease indicators
- Indicators of health disparities
- Chronic disease risk factors (eg, smoking)
- Syndromic surveillance
- Environmental exposures

Number of cities implementing

Abbreviations: HIE, health information exchange; WIC, Supplemental Nutrition Program for Women, Infants, and Children.
Respondent perspectives on health IT and informatics needs and achievements: Interview results

Local sources of data

Interviews with 45 leaders of BCHC LHDs revealed that health departments collect data from a variety of local sources outside the health department to augment their local-level knowledge. Local sources of data included hospital discharge information, emergency medical services (EMS) data, school system data, and data from other city agencies. Thirteen of 16 participating BCHC departments received a variety of data from local hospitals or EMS, including hospital discharge data. One department collected EMS data to track bicycle and pedestrian injuries, homicides, intentional injuries, and narcotics-related information. In addition, some health departments create surveys and other data collection methods as ways to collect the local information they need.

In efforts to build a more comprehensive picture of health citywide, 7 departments reported collecting data from other city agencies, including departments of planning, housing, transportation, education, and law enforcement. Direct data-sharing agreements were also in place between 6 health departments and school systems within their jurisdictions, which allowed access to data on childhood body mass index. Ten health departments reported fielding their own community health surveys, paying for oversampling in their cities on state-administered surveys, such as the BRFSS, or collecting community data in some other way. All of these efforts were focused on gathering data that would allow analyses at small area levels.

Achievements using data

Respondents were asked to highlight their greatest public health achievements using local data. They stated that efforts to collect more data from more sources at increasingly local levels are needed to ensure that the right information is available to guide their efforts to promote and protect health (Table 1).

Most commonly, leaders said that their best achievements were being able to use data to inform policy decisions and using neighborhood-level data to educate local decision makers. They enumerated efforts either to use data to inform internal policy or to provide data to other health departments or community groups in efforts to improve policy. For instance, participants used:

- measures of blood cotinine in the city’s nonsmokers to advocate for smoke-free public spaces;
- dynamic modeling of influenza outbreaks to recommend policies that limited the spread of influenza in schools;
- health disparity data as drivers of overall population health status to inform policy makers on how to address disparities;
- surveillance data to demonstrate the effectiveness of measures implemented to curb childhood obesity; and
- alcohol outlet density data to support community organizations engaged in policy change.

Departments also sought to make public health problems more understandable to decision makers by using neighborhood-level data to highlight specific problems. One department created district-level reports for city council members on topics such as
### TABLE 1 • Achievements Using Data

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<td>Use data to inform policy ( (n = 19) )</td>
<td>“I think what I am most proud of is that here in [City], we really use data to inform programs and policies and for me, that’s very exciting. We’re very much an evidence-based health department. Our commissioner is himself an epidemiologist who understands and loves data. The former commissioner was too. Our mayor is someone who loves data, loves to talk about it. And we really do use local data to inform policy . . . . We have many instances of that, whether it be exposure to mercury in fish that has driven public campaigns targeting communities that have a lot of fish consumption, etc. . . . Diabetes, when we found that nearly a third of adult [city residents] with diabetes were undiagnosed and that really prompted us to, again, improve programs and policies around diabetes and it actually helped launch the hemoglobin A(_{1c}) registry. So I think what I’m most proud of is that we are using data to inform policy and programs.” (Science director)</td>
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<td>Use neighborhood-level data to influence decision makers ( (n = 16) )</td>
<td>“[W]e were able to take the data and then produce council district reports on the data—so numbers broken into 10 different council, no maybe 11 different council districts, and there’s a council person from each district. And so you could compare the data, like what are teen birth rates in this district versus teen birth rates in this district. And so it makes for a great eye-opener and got attention with City Council and just brought a little more political weight to the data, so people could say, ‘Oh, look! People from my community really are heavy smokers compared to everywhere else in the town.’” (Policy director)</td>
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<td>Collaborate to create databases ( (n = 7) )</td>
<td>“I think the collaboration that we created between the health department and the five major academic centers as well as the philanthropy community to develop this database has been probably one of the things that I’ve been the most proud about.” (Lead health officer)</td>
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<td>Merge data sources ( (n = 8) )</td>
<td>“I’d say the second thing I’m most proud of is recently linking some of our STD and HIV/AIDS data. You’d think that those data are often looked at together, but in [City] they haven’t been, and I think in many cities, they’re often very separate sets of services often managed by different divisions or units within the health department. So being able to look at how the risk for a new HIV/AIDS diagnosis increases with each prior STD diagnosis, it’s almost looking at the dose-response of STD infections for future HIV/AIDS infection. One, it’s just a good exercise for us to combine those two sets of data. But two, it really gave us some new insights into how to do prevention around HIV/AIDS within our STD prevention screening and treatment work.” (Policy director)</td>
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<td>Collaborate to access to more data ( (n = 5) )</td>
<td>“I would think that the most recent success is one of our biggest: Is that we have . . . gotten access to [schools’] student health records. And we’re able to look at the BMI data on about 88 000 students and issue a report, which I believe is the largest analysis of childhood obesity in [City] that’s ever been conducted, and really kind of get a handle on that. And then we can use that information—and we have used it—to target our CTG that we got with the schools. I think that’s pretty huge. It’s also, I think, opened the door for other data-sharing with the schools.” (Policy director)</td>
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<td>Use social media and tools ( (n = 2) )</td>
<td>“I’ve also been excited and happy about the use of information that’s out on the social media, the health drive innovation—so a food poisoning app that that’s now getting a lot of coverage actually today. About how do we use data on Twitter to help us drive our food protection efforts, I think has been something I’m proud of.” (Lead health officer)</td>
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Abbreviations: BMI, body mass index; CTG, Community Transformation Grant; STD, sexually transmitted disease.

smoking or teen birth rates, information that became an “eye-opener” for council members whose own districts fared poorly. Another department used data on community-level environmental impacts to make the case for environmental justice regulation, whereas another used data on disparities to spur capital investment in new facilities to provide integrated health and human services.

In addition to using granular data to support arguments for policy change and action, respondents said that departments also used novel combinations of data sets to shed new light on problems. For example, one department noted the value of linking identifiers between sexually transmitted disease and HIV/AIDS data, which are often collected separately by isolated programs. Participants said that they also had
successes with efforts that involved collaborating on data with other agencies and community partners to achieve common goals. As noted previously, some departments worked with schools as a way of accessing childhood body mass index data. Collaboration with schools was the most common, although leaders from one department reported collaborating with their city’s housing authority to track the health of public housing residents as compared with non-public housing residents over time. In addition to forming partnerships to access key data, departments formed partnerships to share data, for example, working with outside entities to create a clearinghouse for data from both public health and nonpublic health sources. Health departments also mined “big data,” including the analysis of social media, such as Twitter.

Data needs

Despite big city health departments’ perceived successes in creating ad hoc sources for local data and effecting policy change and other interventions by using that information, participants uniformly discussed persisting needs for more and higher-quality data (summarized in Table 2). Data specific to smaller local areas were cited as a top need by participants from 14 of 16 departments. Participants also said that they would benefit from access to community-level economic development data, along with more data from school systems, and improved “chronic disease and behavioral risk factor surveillance systems at a small-area level.” Respondents said that very granular data were most valuable for local decision making, whether those decisions were made within health departments or at levels that affected wider city or county policies, as indicated by comments in Table 2.

Participants expressed serious concern about the frequent long delays in getting data from sources such as state health departments, noting that they have sometimes had to wait 2 to 4 years to receive vital statistics data on their jurisdiction. Department heads noted that it was difficult to argue that a certain issue was important to address quickly when they only had outdated data to support their claims. Although some leaders said that they have found inventive ways to get information needed to take action, most said that they remained frustrated with the slow pace and bureaucratic hurdles in getting access to needed data.

Participants believed that their LHDs would benefit from additional data from hospital and ambulatory care settings, particularly data from EHRs. Respondents said that EHRs held significant potential for LHD surveillance, especially for chronic disease monitoring to guide action and geographic “hot spotting” of both communicable and chronic diseases not included in statutory reporting requirements. Some participants also noted that information about consumer perceptions and attitudes would be helpful in certain decision-making processes. For example, one participant explained how information about how far community members would travel to access a local swimming pool would help inform decisions about whether closing certain pools would have adverse effects on surrounding communities. Health department leaders also said that they needed data that would facilitate comparison of health indicators between jurisdictions to determine how their community’s health status compared with others.

**Discussion**

A useful way to view these findings is in the context of the timeliness and geographic specificity of public health data as presented in Figure 4. In the early 1900s, health departments used current and address-level data (cell 3 in Figure 4) to study, understand, and address their problems controlling a limited number of communicable diseases. Public health departments were able to address problems effectively because they had the right information available to them. Timely local data are still the core ingredient for solutions to public health problems. However, as uniformly described by participants in our study, local public health professionals have limited access to geographically specific and timely local data, especially related to chronic disease.

Today, chronic disease surveillance focuses on collecting data at national and state levels. These surveys, including the BRFSS, the Youth Risk Behavior Survey, the Pregnancy Assessment and Monitoring System, the National Health and Nutrition Examination Survey, and the National Health Interview Survey, rarely provide any data below the county level and can take years to be released when their potential benefit to strategic planning is more limited (cell 2 in Figure 4). While these surveillance strategies have developed, few mechanisms have been developed that provided the geographic precision and clinical accuracy necessary to monitor chronic conditions at the local level. Thus, LHDs have lost much of their ability to accurately monitor the health status of their communities with geographically specific and timely data. The chronic disease data available to LHDs do not have the precision to identify health problems at census tract, neighborhood, or even zip code levels; to provide the necessary evidence base to support programmatic and policy solutions; or to measure the impact of interventions. Even when data are available at an address level, as is the situation with mortality data, the delay in their
TABLE 2  Top Data Needs

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<td>More local data (28)</td>
<td>“You know, I think BRFSS is great but, again, it’s not granular enough. It would be really helpful to have something that would give us some reasonable sample sizes at a more granular level. And I know this is always what people say. States don’t care about national data; the counties don’t care about state data. [laughs] And your local communities in the county don’t care about county data. So I think it would behoove us to have something that we could make a little more granular in terms of survey input, assessing people’s behaviors and things like that.” (Science director)</td>
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<td>Claims, electronic health record, hospital data (11)</td>
<td>“I think in an ideal world, we would be able to conduct a local health and nutrition examination survey every three years, but we can’t and most of that is due to resources, both labor and money, and so that’s challenging. That’s really one of the reasons we’re looking at electronic health record surveillance because we’re hoping that that can fill some of the gaps we have.” (Science director)</td>
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<td>Timely data (6)</td>
<td>“I mean, our biggest struggle is having up-to-date data, really. It’s very hard to say, ‘This is a pressing issue’ if your data is from 2008, 2009. Years ago. You know it’s really hard to make people believe you. [laughs] To believe that, it’s still a problem especially if they’re not inclined to believe you in the first place. So that’s our challenge almost across the board, really, in terms of trying to—particularly with mortality data. We just don’t get it in a reasonable amount of time. That’s our biggest struggle is really trying to have the most up-to-date information.” (Policy director)</td>
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<td>Consumer perceptions and attitudes (4)</td>
<td>“I think it would be great to have better data on what people understand about various services that are available and what are their cutoffs or barriers for using those. So for example, we’re dealing with some issues related to community pools—swimming pools—and how to decide which ones of them to keep and which ones to close, and that kind of thing. And we don’t have a really good sense of a resident of a . . . city neighborhood, how far would they be willing to walk; how long would they be able to sit on a bus to go to a pool; or what kind of program that would be attractive to them.” (Science director)</td>
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<td>Comparative data (4)</td>
<td>“We don’t have good comparative data from other cities . . . . We can’t look across jurisdictions at rates of cancer. We can’t look how other cities are doing and what the problem profile looks like on obesity or its causes, so we have lots of data that we’ve generated at the local level that we can’t compare to other cities. I’d really like to be able to learn from other cities.” (Local health official)</td>
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Abbreviation: BRFSS, Behavioral Risk Factor Surveillance Survey.

FIGURE 4  Conceptual Model of Public Health Data Needs in Local Health Departments

Availability drastically limits their impact and places LHDs in the position of implementing solutions today for problems documented more than a year ago (cell 4 in Figure 4).

Some cities have implemented local surveys similar to those implemented nationally to gather subcity data (eg, the Los Angeles County Health Survey and the New York City Health and Nutrition Examination Survey). While effective, local surveys can be costly and require significant staff resources. Completion of the 2013 New York City Health and Nutrition Examination Survey, for example, cost more than $4 million and required the efforts of the New York Department of Health and Mental Hygiene, the Hunter College School of Public Health, and several additional partners. Partly because of the cost, this was the first time that this type of survey had been fielded in New York City since...
better and more timely decisions. Gathering techniques to access the information to make health leaders have implemented “guerilla-style” data-gathering techniques to access the information to make better and more timely decisions. For example, the Boston Public Health Commission (BPHC) investigated an increase in opioid overdose deaths. The BPHC first checked mortality data, but the only data available were 4 years old. The emergency department data it could access were 3 years old. However, because the BPHC has authority over Boston’s EMS, it was able to access real-time, address-level EMS data that allowed it to pinpoint sources of the overdose problem and take action. Like the data used in Boston, several study participants noted the value of clinical data to provide geographically specific and timely data on chronic disease (cell 3 in Figure 4). Public health should strive for “gold standard” data collection and research studies whenever or wherever possible. However, this is often infeasible. Our respondents indicated that they frequently encounter situations that require rapid decisions or warrant immediate action. In these situations, public health leaders may need to use data for operational decision making that may not be “gold standard” but, nonetheless, can impact and improve the health of the public.

The major public health problems have changed over the last century—from communicable to chronic disease. The BCHC LHDs stated that to address population health issues, they must reduce the lag in data access. Data should also change from aggregate (above county level) information (cell 2 in Figure 4) to be as “hyperlocal” as possible, perhaps even to the census tract or address level (cell 3 in Figure 4). Although great advances have been made in chronic disease surveillance at the national and state levels, additional funding needs to focus on developing chronic disease and risk factor surveillance capacity at the local level.

The participants in this study noted the opportunity to leverage the national investment in the EHR infrastructure as a strategy to gain access to chronic disease data that are more timely and geographically specific (cell 3, Figure 4). In the next decade, the total cost of the federal EHR incentive program will exceed $19 billion. Given this investment, determining how these data can be augmented with other contextualizing data (eg, location of fast food outlets, availability of parks, incidents of violent crime, location of public housing, or housing foreclosures), and their use in developing policies with more timely and detailed information should be a priority. Some large urban health departments have already initiated projects that can capitalize on EHR data for public health purposes. For example, New York City is developing the NYC Macroscope as a way to use primary care practice EHRs to track conditions that are important to public health. Work is underway to compare the information in the NYC Macroscope against a gold standard, population-based examination survey to validate the effectiveness of using an EHR as a population health surveillance tool. The public health leaders who participated in this study supported the use of electronic health data to fill the many gaps in local health information. However, clear strategies and “how-to” guides are needed to facilitate partnerships between health care providers and public health leaders to maximize the potential impact of EHRs on population health.

● Conclusion

Monitoring the status of community health is a core function of all public health departments. Public health professionals must have access to current local data on both risk factors and health status to effectively target interventions, wisely allocate resources, and assess the effects of interventions. While the innovations in our largest cities have allowed access to some local data related to chronic disease, LHDs still urgently need timelier and geographically specific data to efficiently and effectively address the most pressing problems in public health.

REFERENCES


