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Gastrointestinal Disease Outbreak Detection Using Multiple Data Streams from Electronic Medical Records

Sharon K. Greene,¹ Jie Huang,² Allyson M. Abrams,¹ Debra Gilliss,³ Mary Reed,² Richard Platt,¹ Susan S. Huang,⁴ and Martin Kulldorff¹

Abstract

Background: Passive reporting and laboratory testing delays may limit gastrointestinal (GI) disease outbreak detection. Healthcare systems routinely collect clinical data in electronic medical records (EMRs) that could be used for surveillance. This study's primary objective was to identify data streams from EMRs that may perform well for GI outbreak detection. Methods: Zip code-specific daily episode counts in 2009 were generated for 22 syndromic and laboratory-based data streams from Kaiser Permanente Northern California EMRs, covering 3.3 million members. Data streams included outpatient and inpatient diagnosis codes, antidiarrheal medication dispensings, stool culture orders, and positive microbiology tests for six GI pathogens. Prospective daily surveillance was mimicked using the space-time permutation scan statistic in single and multi-stream analyses, and space-time clusters were identified. Serotype relatedness was assessed for isolates in two Salmonella clusters. **Results:** Potential outbreaks included a cluster of 18 stool cultures ordered over 5 days in one zip code and a Salmonella cluster in three zip codes over 9 days, in which at least five of six cases had the same rare serotype. In all, 28 potential outbreaks were identified using single stream analyses, with signals in outpatient diagnosis codes most common. Multi-stream analyses identified additional potential outbreaks and in one example, improved the timeliness of detection. Conclusions: GI disease-related data streams can be used to identify potential outbreaks when generated from EMRs with extensive regional coverage. This process can supplement traditional GI outbreak reports to health departments, which frequently consist of outbreaks in well-defined settings (e.g., day care centers and restaurants) with no laboratory-confirmed pathogen. Data streams most promising for surveillance included microbiology test results, stool culture orders, and outpatient diagnoses. In particular, clusters of microbiology tests positive for specific pathogens could be identified in EMRs and used to prioritize further testing at state health departments, potentially improving outbreak detection.

Introduction

IN 2007, FOODBORNE DISEASE OUTBREAKS were associated with over 21,000 reported illnesses in the United States (CDC, 2010b). Health departments (HDs) are commonly notified of focal (e.g., restaurant-associated) outbreaks by passive reports from clinicians or patients. These reports may be incomplete, delayed, and/or non-representative. Outbreaks of intermediate scope, such as those caused by contaminated commercial products, are often detected via laboratory testing. HDs can monitor for unusual increases in passive laboratory-based reports of notifiable diseases (CDC, 2009b) or for clusters of isolates with identical pulsed-field gel electrophoresis (PFGE) patterns (Swaminathan *et al.*, 2001; Gerner-Smidt *et al.*, 2006). However, some gastrointestinal (GI) pathogens are not nationally notifiable (e.g., norovirus, campylobacteriosis), and due to resource limitations at HDs, laboratory testing may be delayed. Generalized outbreaks (e.g., seasonal rotavirus increases) are seldom reported to HDs.

In electronic medical records (EMRs), healthcare systems routinely collect GI disease-related clinical and laboratory data. Using these data may improve the timeliness and representativeness of outbreak surveillance. A prior evaluation of 1 year of EMR data in four states for nine syndromes, including upper and lower GI, did not identify any clusters of public health interest (Yih *et al.*, 2010); however, only

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ambulatory care (AC) diagnoses were used, though other data streams may be more informative. It is also possible that no single clinical data stream is optimal for surveillance, but that multiple syndromic and laboratory data sources could be useful to identify outbreaks.

Space-time clusters in laboratory-based data have been identified for Escherichia coli O157 (Pearl et al., 2006) and Shigella (Jones et al., 2006; Stelling et al., 2009). Syndromic surveillance for GI outbreaks has been evaluated using AC diagnoses (Yih et al., 2005, 2010), emergency department (ED) chief complaints (Balter et al., 2005; Kulldorff et al., 2005), telephone helplines (Caudle et al., 2009), and medication sales (Kirian and Weintraub, 2010; Pelat et al., 2010). To facilitate capacity planning, one hospital conducted univariate temporal analyses of three streams and determined that GI complaints in free text were more useful for rotavirus outbreak detection than either rotavirus antigen laboratory tests or diarrheal disease discharge diagnoses (Levin and Raman, 2005). To our knowledge, no study has used the same patient population to evaluate multiple EMR data streams for spatiotemporal GI outbreak detection.

Prior work using EMR data to detect localized excess influenza-like illness (ILI) suggested that AC diagnoses, reverse transcription-polymerase chain reaction (RT-PCR) tests ordered, and antiviral dispensings were most useful for surveillance (Greene et al., 2011). However, the analogous data streams may not be useful for GI outbreak detection. ILI is caused by respiratory viruses transmitted person-to-person, peaks each winter in temperate climates, and has widespread illness activity. In contrast, GI illness has heterogeneous viral, bacterial, parasitic, and chemical etiologies; is transmitted person-to-person via the fecal-oral route or by a contaminated vehicle; may or may not have a demonstrable seasonal pattern; and the scale may be attributable to a highly localized point source or an internationally disseminated product. True increased ILI activity should be reflected across all streams around the same time, but true increased GI activity may be detectable in only some streams. For example, lower GI syndrome and positive Shigella tests may reflect a shigellosis outbreak, while upper GI syndrome and positive norovirus tests may reflect a norovirus outbreak.

The objectives of this study were to use data from a comprehensive regional health system to (1) create syndromic and laboratory-based data streams for GI disease from EMRs, (2) mimic near real-time prospective surveillance to identify which streams perform well for outbreak detection, and (3) compare results of single stream and multi-stream analyses.

Methods

Study population

Kaiser Permanente Northern California (KPNC) is a large, integrated healthcare delivery system utilizing comprehensive EMRs with inpatient, outpatient, laboratory, and pharmacy data. As of January 2009, KPNC included 18 medical centers and 3.3 million members, representing approximately 22% of the total population residing in 46 counties (951 zip codes) in the Central Valley and San Francisco Bay area. KPNC laboratory-based reports of notifiable diseases are sent to the local HD, which in turn report them to the California Department of Public Health (CDPH) (Backer *et al.*, 2001; California Code of Regulations, 2009).

Data streams

Twenty-two data streams were analyzed, based on syndromic definitions (n=14), prescription drug dispensings (n=1), microbiology tests ordered (n=1), and microbiology test results (n=6) (Table 1). Syndromic definitions were adapted from lists previously developed by a Centers for Disease Control and Prevention/Department of Defense working group (CDC, 2003). Upper GI (UGI) captured vomiting, and lower GI (LGI) captured diarrhea and gastroenteritis (Table 2). Each stream consisted of residential zip code-specific daily episode counts in 2009. Within each stream, an "episode" was the first patient encounter after at least 42 days with no encounter (Jung *et al.*, 2009).

Additional streams were generated but ultimately excluded from analyses. Pathogen-specific ICD-9 codes were excluded because of unreliability, e.g., there were fewer ICD-9 code episodes for *Salmonella* (003.0, 003.20, 003.29, 003.8, 003.9) than laboratory tests positive for *Salmonella*, and ICD-9 codes are assigned before test results become available. Microbiology test results for specific pathogens (e.g., rotavirus antigen tests) were excluded due to low usage.

Univariate single stream analyses

Near real-time prospective surveillance was mimicked by analyzing data "each day." The prospective space-time permutation scan statistic (Kulldorff et al., 2005) was used to detect and evaluate the strength of potential outbreaks. Using a variable-sized cylinder, where the circular base represents space and the height represents time, the method scans the geographic area for potential outbreaks at different locations, with different radii and lengths of time. For each location and cylinder size, a likelihood ratio-based test statistic compares the observed number of cases within the cylinders with what would be expected if the spatial and temporal locations of all cases were independent of each other so that there is no spacetime interaction. As such, it adjusts for any purely spatial and any purely temporal clusters. The cylinder with the maximum likelihood ratio is the most likely cluster, the least likely to have occurred by chance. Calculations were done using SaTScan[™] (www.satscan.org).

The maximum geographical size of the cylinder was set to contain at most 50% of the observed episodes, and the maximum temporal size was set to 14 days. Since the weekly pattern of health-seeking behavior may vary geographically, we adjusted for space by day-of-week interaction, with holidays treated as Sundays and the day after holidays treated as Mondays. The surveillance period was January 1 to December 31, 2009. A 365-day rolling control period established local baselines for each zip code.

To determine statistical significance, 9,999 Monte Carlo simulations (Dwass, 1957) were performed "each day." The recurrence interval (RI) for each cluster represents the length of follow-up required to expect one cluster at least as unusual as the observed cluster by chance (Kleinman *et al.*, 2004). The single stream RIs were statistically adjusted for multiple testing in terms of the thousands of cylinders considered for each data stream, but not for the fact that 22 different single streams were analyzed. In Table 7 below, where only five streams were considered, we also present RIs that were not only adjusted for the many cylinders, but also for the five streams analyzed. This was done by dividing each RI by five.

MULTI-STREAM GASTROINTESTINAL OUTBREAK DETECTION

| Category | # | Data stream | Notes |
|----------------------------------------------|--------|------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| UGI and LGI, ICD-9 | 1 | UGI in AC | ICD-9 codes in Table 2 |
| code based | 2 | LGI in AC | |
| | 3 | UGI in ED | |
| | 4 | LGI in ED | |
| | 5 | UGI hospital discharge | Primary discharge diagnosis |
| | 6 | LGI hospital discharge | |
| | 7 | GI in AC, <5 year-olds | UGI and LGI diagnosis codes combined. |
| | 8 9 | GI in ED, <5 year-olds GI hospital discharge, <5 year-olds | <5 year-olds are the age category at highest risk for rotavirus gastroenteritis (Peck and Bresee 2006), and a population at higher risk for outbreaks in daycare centers and petting zoos. |
| | 10 | UGI in AC with Rx | Antibiotics (Rx) were identified using National |
| | 11 | LGI in AC with Rx | Drug Codes for macrolides, quinolones, |
| | 12 | UGI in ED with Rx | metronidazole, and trimethoprim- |
| | 13 | LGI in ED with Rx | sulfamethoxazole |
| | 14 | GI hospital admission | Text strings for vomiting, diarrhea, and gastroenteritis, with exclusions for chronic diarrhea, pregnancy, chemotherapy, and intoxication (Chapman <i>et al.</i> , 2010; Chapman 2011) |
| Prescription antidiarrheals | 15 | Antidiarrheal dispensing | NDCs for diphenoxylate and loperamide. Note that a portion of these dispensings would reflect prescriptions in advance of international travel, rather than acute illness treatment. Over-the-counter anti-diarrheal agents were not considered, as relevant clinician recommendations for symptomatic treatment would be captured only sporadically in EMR text fields. |
| Microbiology tests ordered in any setting | 16 | Stool culture tests | |
| Microbiology tests | 17 | Campylobacter | Positive stool culture test. Analyzed according to |
| positive for GI | 18 | Salmonella | date test ordered, not according to lagged date |
| pathogens | 19 | Shigella | when test results became available. |
| 1 0 | 20 | E. coli O157:H7 | |
| | 21 | Vibrio parahaemolyticus | |
| | 22 | Cryptosporidium | Positive stain |

TABLE 1. DATA STREAM DEFINITIONS

AC, ambulatory care; ED, emergency department; GI, gastrointestinal; LGI, lower gastrointestinal; Rx, antibiotic prescription; UGI, upper gastrointestinal.

For streams other than tests positive for GI pathogens, all clusters from prospective analyses with RI of > 365 days were identified. Clusters from the same stream on consecutive days and overlapping in space were grouped together into "cluster sequences." These cluster sequences were then compared across streams, and sequences with at least 1-day temporal overlap and spatial overlap (cluster center in other cluster) were further grouped together into "potential outbreaks."

For the six streams of tests positive for GI pathogens, clusters with RI of >60 days were identified. This lower RI threshold was selected because of the sparseness of microbiology data, and because the trigger to begin a cluster investigation may be lower for these streams, which are more specific for acute infections than the syndromic streams.

The epidemiological interpretation of clusters is subjective. Clusters may be prioritized for possible investigation by simultaneously weighing (1) the number of observed cases (the greater, the more urgent the need for a public health intervention), (2) the observed/expected number of cases (the greater, the more excess risk), (3) the RI (the greater, the less likely the observed clustering is due to chance), and (4) the degree of localization (a smaller radius more strongly suggests a common source or localized person-to-person transmission).

Multi-stream analyses

Multivariate analyses use multiple streams in the same statistical analysis (Burkom *et al.*, 2005; Kulldorff *et al.*, 2007; Rolka *et al.*, 2007). Five streams were included in multivariate analyses: three microbiology-based streams were selected because cluster detection could prompt case interviews and PFGE testing of isolates, and two syndromic streams were selected based on frequency of episodes and contribution to potential outbreak detection in single stream analyses. To avoid multicollinearity across streams, a patient appearing in more than one stream within 14 days was retained in only one stream. A priority order of increasing frequency was used so

| Syndrome | Definition | International Classification of Diseases, Ninth Revision codes |
|-------------|------------------------------------------------------------------------------|-------------------------------------------------------------------------|
| UGI illness | Epidemic vomiting syndrome | 078.82 |
| | Gastritis and duodenitis | 535.0, 535.4, 535.5, 535.6 |
| | Persistent vomiting Nausea and vomiting | 536.2 787.0 |
| LGI illness | Other bacterial food poisoning | 005.89, 005.9 |
| | Intestinal infection due to other organisms | 008.49 |
| | Bacterial enteritis unspecified | 008.5 |
| | Enteritis due to other viral enteritis | 008.69 |
| | Intestinal infection due to other organism not elsewhere classified | 008.8 |
| | Ill-defined intestinal infections | 009 |
| | Regional enteritis | 555.0, 555.1, 555.2 |
| | Other and unspecified noninfectious gastroenteritis and colitis | 558.2, 558.9 |
| | Unspecified disorder of intestine | 569.9 |
| | Visible peristalsis | 787.4 |
| | Diarrhea | 787.91 |

TABLE 2. SYNDROMIC DEFINITIONS FOR UPPER AND LOWER GASTROINTESTINAL ILLNESS

LGI, lower gastrointestinal; UGI, upper gastrointestinal.

that patients would be retained in the stream in which they were most proportionally informative: *E. coli* O157:H7, *Salmonella*, *Campylobacter*, LGI in ED, and LGI in AC.

With these five streams, the multivariate scan statistic simultaneously searched for clusters in all 31 possible combinations of one or more streams, adjusting for the multiple testing inherent in both the thousands of cylinders evaluated for each combination and in the 31 data stream combinations evaluated. On each surveillance day, the combined log-likelihood was defined as the sum of the individual log-likelihoods for those streams with more observed events than expected (Kulldorff *et al.*, 2007). The maximum cylinder size and other settings were the same as for the single stream analyses.

State HD data

For context, but not for direct comparison with potential outbreaks identified using KPNC data, we compiled a list of GI disease outbreaks known to CDPH occurring in non-institutional settings affecting any of the 16 counties for which KPNC had $\geq 10\%$ population coverage. One author (D.G.) provided preliminary GI illness outbreak reports, which had been sent to CDPH by local HDs soon after outbreak detection. CDPH's final foodborne disease outbreak reports reflected the best available information after completed outbreak investigations. The preliminary and final outbreak reports were unlinked and represented separate information sources.

The CDPH state laboratory performs *Salmonella* serotyping. Previously collected serotype information was obtained to evaluate the two potential *Salmonella* outbreaks with the most observed cases identified in KPNC EMR data.

Results

Figure 1 shows the frequencies and seasonal patterning of the 22 data streams. The relative frequencies of pathogens (Fig. 1A) were consistent with the most common laboratoryconfirmed infections in 2009 at the California FoodNet site (CDC, 2010a).

Single stream analyses

In an illustrative potential outbreak, a signal occurred on November 9, 2009 for stool culture tests ordered, with signals continuing over the subsequent four days (Tables 3 and 4, #19). Ultimately, 18 tests were ordered in one zip code over 5 days, with less than four tests expected. Statistically, this was very unlikely to occur by chance. No corresponding cluster was detected of tests positive for any of the six specific pathogens under surveillance, but the cluster could have reflected an outbreak of a viral pathogen.

In single stream analyses of 16 non-pathogen-specific streams, 24 potential outbreaks were detected using a 365-day RI threshold (Table 4). Of the 16 streams, two had no clusters with RI of > 365 days: UGI with Rx, in the AC and ED settings. Three streams each contributed to the identification of >5 potential outbreaks, all diagnoses in outpatient settings: LGI in ED, UGI in ED, and LGI in AC.

In single stream analyses of six pathogen-specific streams, five potential outbreaks (two *Campylobacter* and three *Salmonella*) were detected using a 60-day RI threshold (Table 5). In potential outbreak B, serotype information was available for five of the six isolates, all from patients residing in one zip code. All isolates were *Salmonella enterica* serotype Thompson. Nationwide, *Salmonella* serotype Thompson represents only 1% of all serotyped *Salmonella* isolates (CDC, 2008), which suggests this may have been an outbreak with a common source or person-to-person transmission. This event was only detected using microbiology data, not with any of the less specific and noisier syndromic streams.

In potential outbreak C, serotype information was available for all 10 isolates: four Enteritidis (the most common serotype (CDC 2010a)), three Montevideo, one Heidelberg, one Infantis, and one Paratyphi B L(+) tartrate +. This mix of serotypes over 10 days across 71 zip codes is unlikely to represent a common source outbreak.

Multi-stream analyses

Multivariate analyses detected six potential multi-stream outbreaks (Table 6), two of which were not detected in single stream analyses. Some potential outbreaks were detected by both single and multi-stream analyses, but at different RI strengths. Table 7 shows examples where multi-stream

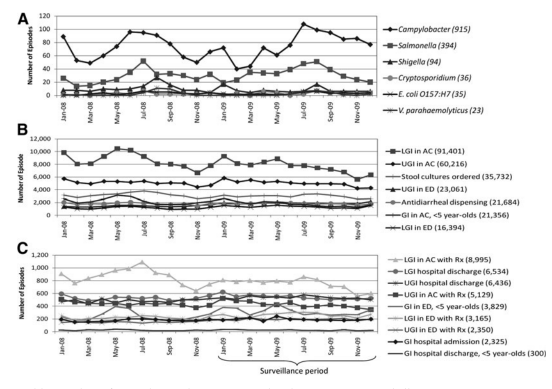


FIG. 1. Monthly number of episodes in data streams related to gastrointestinal illness in Kaiser Permanente Northern California, 2008–2009. **(A)** Data streams for microbiology tests positive for specific pathogens (n=6). **(B)** Non-laboratory-based data streams with \geq 10,000 episodes in 2009 (n=7). **(C)** Non-laboratory-based data streams with <10,000 episodes in 2009 (n=9). Legend indicates data stream and number of episodes in 2009.

analyses detected a cluster with a higher RI than single stream analyses, and vice versa. In one example centered in Firebaugh over 6 days, multi-stream analysis resulted in more timely detection of a potential outbreak than single stream analysis (Fig. 2). Maps of clusters on all days in 2009 with signals in either single or multi-stream analyses are available (see Supplementary Material available online at www.liebertonline.com/fpd).

Outbreaks reported to CDPH

Twenty-four GI outbreaks affecting the study area in 2009 were reported to CDPH. Three were multi-state outbreaks associated with contaminated commercial food products (Nielsen *et al.*, 2010; CDC, 2009a; FDA, 2009). Two additional outbreaks were laboratory-confirmed: a restaurant-based norovirus outbreak (n=8) and a home-based *E. coli* O157:H7 outbreak (n=5). The remaining 19 reported outbreaks had no laboratory-confirmed etiology, were mostly in restaurants

and child day care centers, and had few reported cases (median, 10; range, 3–35).

Discussion

GI disease-related data streams can be generated from EMRs and prospectively used to identify potential outbreaks in settings with extensive regional coverage, such as KPNC. Different surveillance systems will have different strengths and weaknesses, and one should not expect EMR data streams to detect all outbreaks reported to CDPH and vice versa. Not all true outbreaks are recognized, and not all recognized outbreaks are reported (CDC, 2010b). In addition, the three multi-state outbreaks known to CDPH may have clustered only in time but not in space within the catchment area, so that they could not be detected by spatial methods. Also, with a median number of nine cases, the other 21 outbreaks reported to CDPH would have been difficult to detect. Given

 Table 3. Illustrative Potential Outbreak: Stool Culture Tests Ordered over Five Consecutive Days in One Zip Code

| Start date | Signal date | Number of zip codes | Observed (O) | Expected (E) | O/E | Recurrence interval (days) |
|------------|-------------|------------------------|--------------|--------------|------|-------------------------------|
| 11/9/09 | 11/9/09 | 1 | 9 | 0.6 | 14.4 | 3,333 |
| 11/9/09 | 11/10/09 | 1 | 11 | 1.4 | 8.1 | 769 |
| 11/9/09 | 11/11/09 | 1 | 14 | 2.1 | 6.8 | 2,000 |
| 11/9/09 | 11/12/09 | 1 | 17 | 2.7 | 6.4 | 5,000 |
| 11/9/09 | 11/13/09 | 1 | 18 | 3.4 | 5.4 | 10,000 |

| | | | | Division of KDNIC | | L | Details as of date with maximum recurrence interval | ith maximun | і геситепсе | interval | |
|-----------------------|------------------------------|----------------|------------------------|----------------------------------|----------------------------------------------|-----------------------|-----------------------------------------------------|-----------------|-----------------|-------------|------------------------|
| Potential outbreak | Central location | Radius (km) | Number of zip codes | population in geographic area | Data stream | Signal date (2009) | Number of days in cluster | Observed (O) | Expected (E) | O/E | Recurrence interval |
| 1 | San Francisco | 14 | 64 | 7.8 | GI in AC, <5 year-olds | 1-Jan | 14 | 88 | 46.5 | 1.9 | 5,000 |
| | Sausalito | 18 | 70 | 6.5 | LGI in AC | 1-Jan | 14 | 246 | 161.5 | 1.5 | 10,000 |
| 7 | San Jose | 12 | 49 | 6.2 | UGI in ED | 4-Jan | 14 | 48 | 20.8 | 2.3 | 3,333 |
| ი. ე | Westley | 36 | 34 | 10.1 | LGI in ED | 23-Jan | 80 j | 39 | 15.4 | 2.5 | 769 |
| 4 | Half Moon Bay | 29 | 67 | 6.9 | LGI in AC | 27-Jan | 13 | 563 | 435.9 | 1.3 | 10,000 |
| | Sausalito San Francisco | 24 1 | 102 6 | 11.3 03 | GI in AC, <5 year-olds I CI in AC with Rv | 14-Feb 16-Feb | 14 | 159 | 103.8 | 1.5 דק | 500 625 |
| IJ | Hilmar | 32 - | 31 | 0.0 | UGI in ED | 15-Feb | 11 | 36 | 12.3 | 2.9 | 5,000 |
| | Crows Landing | 35 | 31 | 2.5 | LGI in ED | 19-Feb | 14 | 34 | 12.5 | 2.7 | 526 |
| , | Huron | 195_{-} | 210 | 12.5 | UGI hospital discharge | 24-Feb | υ | 26 | 7.9 | 3.3 | 2,500 |
| 91 | Sacramento | ► ç | 19 | 2.0 | LGI in ED | 30-Mar | 12 | 31 | 10.2 | 3.0 | 1,111 |
| | Sacramento | 13 | 27 | 5.9 | GI hospital discharge, | 29-Apr | 13 | x | 1.2 | 6.9 | 629 |
| x | Merced | 41 | 24 | 17.0 | LGI in FD | 5-May | 11 | 17 | 1 9 | ц У | 435 |
| D | Firebauch | 101 | 134 134 | 0.41 | | 14-May | 14 | 413 413 | 309.9 |) () (| 10.000 |
| | Chowchilla | 106 | 151 | 6.8 | LGI in ED | 14-Mav | 14 | 63 | 46.9 | 2.0 | 10,000 |
| | Linden | 53 | 89 | 9.6 | UGI hospital discharge | 23-May | 11 | 31 | 11.2 | 2.8 | 417 |
| | Merced | 67 | 138 | 2.9 | GI in AĊ, <5 year-olds | 26-May | ~ | 93 | 48.7 | 1.9 | 3,333 |
| | Dos Palos | 80 | 72 | 13.6 | UGI in ED | 28-May | 14 | 77 | 36.0 | 2.1 | 10,000 |
| | Fowler | 8 | 7 | 0.1 | LGI in AC | 31-May | 1 | IJ | 0.1 | 38.4 | 769 |
| | Valley Springs | 55 | 81 | 6.6 | LGI in AC | 1-Jun | 14 | 324 | 240.5 | 1.4 | 370 |
| | Lodi | 30 | 39 | 11.2 | UGI in ED | 2-Jun | ŝ | 21 | 5.0 | 4.2 | 1,667 |
| 6 | Berkeley | | 18 | 2.3 | LGI in ED with Rx | 21-Jun | 13 | 14 | 2.6 | 5.4 | 714 |
| 10 | Winters | 16 0 | т, т | 0.5 | LGI in ED | 21-Jun | 13 | 15 | 2.9 | 5.1 | 400 |
| 11 | San Jose | ⊃ç | | 4.7 | | 24-Jun | 4 ç | χí | 0.0 | 14.3 | 1,067 |
| | Santa Clara Mountain View | 5 <u>1</u> 0 | 40 23 | υ.υ υ.υ | CI hoenital discharoo | 1-Jul 3-Iul | 12 م | n 0 1 | 28.U | 2.3 10 7 | 1 479 |
| | | | Ì | i | <5 vear-olds | m |) |) | 1.0 | | / / |
| | Santa Clara | 12 | 57 | 7.0 | LGI in ED | 21-Jul | 11 | 57 | 24.0 | 2.4 | 10,000 |
| | San Jose | ~ | 32 | 4.1 | GI in ED, <5 year-olds | 16-Aug | 13 | 16 | 3.2 | 5.0 | 3,333 |
| | Mountain View | 19 | 76 | 5.6 | LGI in ED | 30-Aug | 14 | 78 | 38.1 | 2.1 | 10,000 |
| 12 | Montague | 366 | 257 | 26.0 | LGI in AC | 11-Jul | ŋ | 318 | 225.9 | 1.4 | 10,000 |
| | Kenwood | 16 | 21 | 2.3 | LGI hospital discharge | 17-Jul | 4 | 14 | 2.3 | 6.0 | 1,111 |
| 13 | Campo Seco | 52 | 74 | 5.0 | GI in AC, <5 year-olds | 25-Jul | 13 | 72 | 35.9 | 2.0 | 2,500 |
| 14 | Vallejo | 0 | - | 0.5 | GI in ED, <5 year-olds | 29-Jul | 2 | 4 | 0.1 | 46.7 | 417 |
| 15 | Stockton | 42 | 57 | 7.3 | UGI in AC | 11-Oct | (n) | 57 | 26.1 | 2.2 | 606 |
| 16 | Hidden Valley Lake | 48 | 31 | 0.8 | LGI in AC | 17-Oct | 13 | 120 | 72.1 | 1.7 | 556 |
| 17 | Guinda | χ χ | 54 0 | 1.7 | | 26-Uct | 12 | 53 5 | 22.8 | 5.2 | 3,333 |
| 17 | San Francisco | 7 | V | C.U | Antidiarrheal dispensing | 29-Oct | Ч | ø | 0.0 | 14.4 | 1,111 |

(continued)

| Potential outbreak Centre | | | | Davesutage of VDNIC | | 1 | | | | | |
|------------------------------|------------------|----------------|------------------------|----------------------------------|--------------------------|-----------------------|------------------------------------------------------------------------|-----------------|-----------------|------|------------------------|
| | Central location | Radius (km) | Number of zip codes | population in geographic area | Data stream | Signal date (2009) | Signal date Number of days Observed Expected (2009) in cluster (O) (E) | Observed (O) | Expected (E) | O/E | Recurrence interval |
| 18 Rough a | Rough and Ready | 46 | 43 | 1.9 | LGI in AC with Rx | 2-Nov | 14 | 22 | 5.8 | 3.8 | 714 |
| 19 Elk Ğrove | ve | 0 | 1 | 0.8 | Stool cultures ordered | 13-Nov | ъ | 18 | 3.4 | 5.4 | 10,000 |
| 20 San Rafael | ael | 19 | 42 | 4.1 | GI hospital admission | 18-Nov | 13 | 15 | 2.6 | 5.7 | 10,000 |
| 21 San Jose | Ċ, | 1 | 7 | 0.8 | LGI in AC with Rx | 25-Nov | 9 | 6 | 0.9 | 9.6 | 606 |
| 22 San Francisco | ncisco | ŋ | 32 | 2.7 | Antidiarrheal dispensing | 2-Dec | 4 | 30 | 10.2 | 3.0 | 500 |
| 23 Boulder Creek | Creek | 27 | 60 | 5.3 | LGI in ED | 8-Dec | 9 | 32 | 10.9 | 3.0 | 1,667 |
| 24 Fairfield | 1 | 24 | 11 | 5.9 | UGI in ED | 23-Dec | 13 | 57 | 24.8 | 2.3 | 10,000 |
| Capay | | 34 | 16 | 0.7 | GI in ED, <5 year-olds | 25-Dec | ю | J. | 0.2 | 27.6 | 588 |

TABLE 4. CONTINUED

AC, ambulatory care: ED, emergency department; GL gastrointestinal illness; LGL, lower gastrointestinal illness; Rx, antibiotic prescribed; UGL, upper gastrointestinal illness; KPNC, Kaiser Permanente Northern California.

| Table 5. Clusters from Single Stream Analysis with Recurrence Interval of >60 Days in Data Streams for Microbiology Tests Positive for Gastrointestinal Pathogens |
|-------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|-------------------------------------------------------------------------------------------------------------------------------------------------------------------|

| T CLCCITUAXE OL TV | | | |
|--------------------|----------------------------------|------------------------------------------------------|------------|
| n ea | population in geographic area | Number of population in zip codes geographic area | f popu |
| | 1.0 | 17 1.0 | 3 17 1.0 |
| | 2.1 | 3 2.1 | 5 3 2.1 |
| | 10.7 | 71 10.7 | 27 71 10.7 |
| | 5.9 | 24 5.9 | 21 24 5.9 |
| | 4.8 | 13 4.8 | 23 13 4.8 |

KPNC, Kaiser Permanente Northern California.

| | | | | Ι, | V I. | ſ | 1 1 | 17 17 | | Ċ | | | Ċ | []- | 1~ | | 11111211C_11111TAT | |
|-----------------------------------|----------------------------|--------|------------------|-----|-----------|-------|-----|-----------|---------|-----|---------------|---------|---|------------|------|--------------------|--------------------|--------|
| Dotantial authord | L Control | Dadine | Padine Mumber of | Ĺ | TPI IN AC |) | Ľ | LGI IN EU | | Can | Campylobacter | crer | 3 | Salmonella | 10 | Cional data | Mumber of date | |
| Fotential outorea from Table 4 | | (km) | zip codes | 0 | Е | O/E O | 0 | Е | O/E O E | 0 | | O/E O E | 0 | | O/E | ықпа ише (2009) | (2009) in cluster | RI |
| 1 | Sausalito | 19 | 83 | 327 | 229.3 | 1.4 | 81 | 64.2 | 1.3 | I | I | I | 2 | 0.9 | 2.3 | 1-Jan | 14 | 10,000 |
| 8 | Firebaugh | 101 | 137 | 404 | 332.7 | 1.2 | 82 | 42.1 | 2.0 | 4 | 1.9 | 2.1 | Ļ | 0.6 | 1.6 | 12-May | 14 | 10,000 |
| None | Arnold ^a | 103 | 190 | 693 | 569.7 | 1.2 | 166 | 124.8 | 1.3 | 9 | 5.6 | 1.1 | I | I | I | 15-Jul | 13 | 10,000 |
| 11 ^c | Santa Clara | 12 | 57 | 343 | 331.9 | 1.0 | 99 | 27.7 | 2.4 | I | I | I | I | I | I | 24-Jul | 14 | 10,000 |
| 11 ^c | Sunnyvale | 14 | 62 | 274 | 244.6 | 1.1 | 47 | 18.1 | 2.6 | | 3.8 | 1.9 | 9 | 2.1 | 2.9 | 30-Aug | 13 | 10,000 |
| None | Discovery Bay ^b | 0 | 1 | | 0.5 | 15.0 | I | I | I | I | I | I | Ļ | 0.03 | 36.5 | 26-Oct | 2 | 417 |

AC, ambulatory care; ED, emergency department; LGI, lower gastrointestinal illness; UGI, upper gastrointestinal illness; O, observed; E, expected. Both of these correspond to the same potential outbreak from single stream analysis.

the 22% KPNC population coverage, on average, only two cases in each outbreak would be KPNC members. Thus, monitoring EMR streams would complement rather than replace traditional outbreak detection systems, which typically detect outbreaks in well-defined settings with no laboratoryconfirmed pathogen. Several EMR streams emerged as most promising for GI

outbreak detection. First, microbiology test results are very specific for acute enteric illness, and even with a low RI, a cluster suggestive of a true outbreak was identified of genetically related isolates (Table 5, potential outbreak B). Also, stool culture orders reflect clinician suspicion of acute illness and disproportionately represent patients with bloody diarrhea and diarrhea duration of ≥ 3 days (Scallan et al., 2006); it is unknown whether the intriguing potential outbreak identified (Table 3) reflected a true outbreak or something else, but it represents the type of event that public health officials may be interested in prospectively detecting. Finally, outpatient diagnoses contributed to the most potential outbreaks (Table 4). Data are rapidly available in the KPNC EMR: diagnoses and stool culture orders typically within one day, and positive stool culture test results in a median of three days. Multi-stream analyses could identify potential outbreaks too faint for detection in single stream analyses (Table 7) and improve the timeliness of detection (Fig. 2), but were not consistently superior or inferior to single stream analyses.

Several limitations should be noted. First, analyses were by zip code of patient residence, so point source outbreaks where people congregate, then disperse, may be missed. Second, only a portion of the total population are KPNC members, and only a fraction of members with GI illness will seek care or submit a stool specimen and thus appear in an EMR; generally, about 20% of patients with an acute diarrheal illness seek medical care, and 4% submit a stool specimen (Jones *et al.*, 2007). Hence, an EMR-based surveillance system cannot be expected to detect very small outbreaks. Third, we evaluated only one geographical area during 1 year. The apparent relative strengths across streams could be different in other places or years.

Microbiology test results in EMRs seem to be especially promising for outbreak detection. Given limited resources and competing priorities within HDs, there are delays in serotyping and an inability to perform PFGE testing on all isolates. Currently at CDPH, Salmonella isolates are prioritized for investigation (e.g., patient interview and/or PFGE testing) based on the presence of an unusual serotype, an increase in a common serotype, or recognized clusters. An alert of a cluster of Campylobacter isolates might trigger CDPH to collect case report details or conduct PFGE testing, which are non-routine activities for campylobacteriosis. HDs could strengthen cooperative partnerships with healthcare systems like KPNC, such that in addition to routinely submitting their isolates to the HD for possible further testing, laboratories in these healthcare systems could provide HDs with counts of microbiology tests ordered and positive, by zip code, for automated daily analyses at the HD, in order to identify unusual space-time clustering. In concert with other strategies for near real-time laboratory-based surveillance (Nielsen et al., 2006; Miller et al., 2010), this could more efficiently prioritize testing and patient interviews, potentially improving outbreak detection.

RI

Table 6. Clusters from Multi-Stream Analysis with Recurrence Interval (RI) of >365 Days, as of the Date with the Maximum

MULTI-STREAM GASTROINTESTINAL OUTBREAK DETECTION

| Did single or multi-stream analysis detect cluster with higher recurrence | Signal date | from single st not adjusted fo streams | ce interval ream analysis, or the five data evaluated | from single st adjusted for | ce interval ream analysis, the five data evaluated | Recurrence interval from multi-stream analysis, adjusted for the 31 combinations |
|---------------------------------------------------------------------------------|----------------------------|----------------------------------------------|----------------------------------------------------------------|--------------------------------|-------------------------------------------------------------|----------------------------------------------------------------------------------------|
| interval? | (2009) | LGI in AC | LGI in ED | LGI in AC | LGI in ED | of the five data streams evaluated |
| Multi-stream | 7-May 15-Jul 26-Oct | 33 2,500 227 | 118 189 | 7 500 45 | 24 38 - | 435 10,000 417 |
| Single stream | 30-Mar 10-Jul 21-Jul | 5,000 | 1,111 _ 10,000 | 1,000 | 222 2,000 | 84 313 2,000 |

| Table 7. Si | ix Illustrative | EXAMPLES IN | м Wнісн | Single | AND | Multi-Stream | ANALYSES |
|-------------|-----------------|--------------|-----------|----------|------|--------------|----------|
| | Detect th | ie Same Clus | STER WITH | 1 Diffei | RENT | Strength | |

AC, ambulatory care; ED, emergency department; LGI, lower gastrointestinal illness.

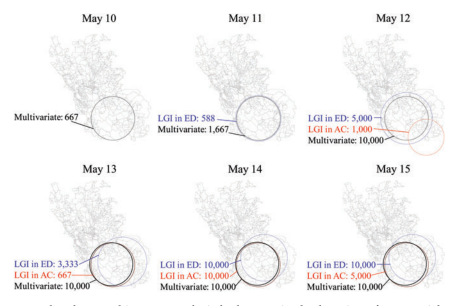


FIG. 2. Illustrative example where multi-stream analysis had more timely detection of a potential outbreak than single stream analysis. On May 10, multivariate analysis first detected a cluster, with a recurrence interval of 667. The next day on May 11, the single stream analysis of lower gastrointestinal illness (LGI) in the emergency department (ED) first detected the same cluster. On May 12, the single stream analysis of lower gastrointestinal illness (LGI) in ambulatory care (AC) detected a cluster, but it was geographically offset. On May 13, the single stream analysis of LGI in AC detected the same cluster that the multivariate analysis first detected 3 days earlier. Color images available online at www.liebertonline.com/fpd

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Disclosure Statement

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