Genetic Diversity as a Marker for Timing Infection in HIV-Infected Patients: Evaluation of a 6-Month Window and Comparison With BED

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Background. It has been reported that the increase in human immunodeficiency virus (HIV) sequence diversity in drug resistance surveillance specimens may be used to classify the duration of HIV infection as <1 or >1 year. We describe a mixed base classifier (MBC) optimized to categorize the duration of subtype B infections as <6 or >6 months on the basis of sequences for drug resistance surveillance specimens and compared MBC findings with those of serologic methods.

Methods. The behavior of the MBC was examined across a range of thresholds for calling mixed bases. MBC performance was then evaluated using either complete pol sequences or sites reflecting evolutionary pressures (HLA selection sites, sites that increased in entropy over the course of infection, and codon positions).

Results. The MBC performance was optimal when secondary peaks on the sequencing chromatogram accounted for at least 15% of the area of primary peaks. A cutoff of <0.45% mixed bases in the pol region best identified recent infections (sensitivity = 82.7%, specificity = 78.8%), with improvement achieved by analyzing only sites that increased in entropy.

Conclusions. In an extended data set of 1354 specimens classified by BED, the optimized MBC performed significantly better than a simple MBC (agreement, 68.98% vs 67.13%). If further validated, the MBC may prove beneficial for detecting recent infection and estimating the incidence of HIV infection.

The early symptoms of human immunodeficiency virus (HIV) infection may not prompt patients to seek medical attention. Consequently, most HIV infections are diagnosed among people who are chronically infected [1], with the date of seroconversion remaining unknown. Nevertheless, correct identification of recent HIV infections (RHIs), defined as infections <6 months old, is critical to public health for several reasons. First, estimates of RHIs in a specified population over a defined period are needed to assess the incidence of HIV infection. This information is necessary to determine trends in the HIV epidemic and to evaluate the success of prevention strategies. Second, contact tracing of RHI patients is critical because it is believed that high rates of onward HIV transmission are associated with elevated viral loads during early infection [2]. Third, because viral strains in RHI patients are closely related to the virus population found in the transmitting partner, identification of RHI offers unique opportunities to improve our understanding of the biology of HIV transmission [3]. However, current methods used to tease apart recent from established infections have severe shortcomings [4].
Incident infections may be identified during the course of routine diagnostic testing. At the simplest level, discordant results of serologic tests (i.e., a negative result followed by a positive result) over a brief and defined period accurately identifies RHI. Alternatively, during the first 2–4 weeks of infection and before seroconversion, the detection of viral capsid (p24) or HIV RNA/DNA in a serology-negative specimen is evidence of RHI [5, 6]. However, these methods will only identify incident infections in the minority of patients who are either frequently tested or who present for care during acute infection.

Beyond the 2–4-week window, HIV-specific immunoglobulin G (IgG) appears as a result of the host adaptive immune response. This appearance is the basis of serologic assays used to identify RHI. The “detuned” approach consists in retesting HIV-positive samples with a less sensitive diagnostic test. Samples negative with the less sensitive assay are classified as RHI [7]. The Calypte BED-CEIA is a capture enzyme immunoassay that estimates the time since seroconversion by measuring the level of HIV-specific IgG relative to the total level of IgG [8]. Alternatively, the avidity of HIV antibodies, a proxy for the maturity of the antibody response, can be used to identify RHI [9]. In Canada, national HIV surveillance relies on the BED assay, which distinguishes infections with a duration of <155 days from those with a duration of >155 days. However, the assay has a number of limitations. First, HIV-specific antibody levels can decrease during infection, either as patients progress to AIDS or when viral load is suppressed [10]. As a result of this decrease, BED misclassifies such infections as recent. Second, the recommended cutoff varies between subtypes, from 155 days for subtype B to 360 days for subtype C [4, 11]. Thus, the application of BED to a population with a multitude of viral subtypes can be challenging.

An alternative approach to determining the stage of infection has been to measure the genetic diversity of HIV within an individual [12]. Previous work showed that sexually transmitted HIV infection is established by only a limited number of viruses [13, 14], with viral genetic diversity increasing subsequently [15, 16]. The proportion of “mixtures” during population-based sequencing, which reflect viral population polymorphisms, may be used as a proxy for within-patient genetic diversity. In their pioneering work, Kouryos et al determined that a mixed-base cutoff of \( Y_{50} = 0.5\% \) achieved a sensitivity of 86.8% and a specificity of 70% in predicting infections with a duration of <1 year [12]. Here, given the existing body of work associated with BED use, we examined whether the classifier would provide concordant classifications at the 155-day BED cutoff. In addition, because the number of mixed bases in a nucleotide sequence implicitly depends on the threshold for calling mixed bases, we tested different thresholds with the classifier. Finally, because selective pressure on the \( \text{pol} \) gene is not evenly distributed, we evaluated whether mixed bases in subsections of \( \text{pol} \) provided greater resolution of recent versus established infections.

**METHODS**

**Study Populations**

The Canadian HIV Strain and Drug Resistance Surveillance Program (SDR) tracks subtype and transmitted drug resistance among patients in Canada who have newly diagnosed HIV infection and are antiretroviral (ART) naive [17]. Remnant HIV diagnostic specimens, along with basic epidemiological data, including exposure category, are sent to the National HIV and Retrovirology Laboratories (NHRL) in Ottawa, Canada, for genotyping. Before specimens are genotyped, they are carefully pedigreed to ensure they originate from the first diagnosis of HIV in that patient. Here, 1450 samples received from Western Canada between 2002 and 2008 were analyzed. The SDR is approved by the Health Canada Research Ethics Board. Additional external sequences originating from patients with RHI were provided by the Jewish General Hospital in Montreal, Canada, the BC Centre for Excellence in Vancouver, Canada, and the National Cancer Institute in Bethesda, Maryland. Sequences from patients with chronic infection were obtained from published work [18]. All sequences originated from ART-naive patients.

**Stage of Infection**

Three data sets were assembled (Supplementary Figure 1), with 2 containing sequences associated with a well-known infection duration and 1 containing sequences associated with an unknown infection duration. Specimens were classified as recent if viral p24 antigen was detected in the sample in the absence of antibodies or if the patient tested negative for HIV within the 155 days prior to diagnosis. Specimens were classified as established if they were collected at least 155 days after the initial HIV diagnosis.

The NHRL training data set consisted of 96 specimens, of which 66 were from recent infections and 30 were from established infections. An additional 237 sequences (96 recent and 141 established) from other laboratories or the literature [18] were added to the NHRL training data set to form the full training data set. In total, the full training data set contained 333 sequences corresponding to 162 recent and 171 established infections.

The NHRL sequences for which the duration of infection was unknown were classified according to results of BED (performed according to manufacturer instructions) as having a duration of <155 days or >155 days [8]. This third independent data set of 1354 specimens (440 recent and 914 established) is denoted hereafter as the BED data set. Other than the 171 sequences from established infections in the training data sets, all samples originated from patients with newly diagnosed infection.
Amplification and Sequencing

Viral RNA was extracted from 0.5-mL serum samples, and pol was reverse transcribed, amplified, and Sanger sequenced as previously described [19]. The lower limit of sensitivity was 1000 copies/mL. Contigs were assembled and aligned to the National Center for Biotechnology Information HIV type 1 reference genome (accession number: NC_001802), using BioEdit 7.0.9 [20]. The obtained pol sequences covered 1305 base pairs, including the protease gene (bases 1–297) and a portion of the reverse transcriptase gene (bases 1–1008). Sequences from samples collected before 2005 were already in GenBank [19] and all new sequences were deposited (accession numbers: HM468499-HM468608, HM468610-HM468617, HM468619-HM468740, HM468750, HM468752-HM468766, HM468768-HM468813, HM468815, HM468817-HM468821, HM468823, HM468825-HM468853, HM468855-HM468870, HM468872, HM468874-HM468875, HM468877-HM468906, HM468908-HM468914, HM468916-HM468936, HM468938-HM468959, HM468964-HM468975, HM468978-HM468982, HM468984-HM468992, HM468994-HM468999, HM469001, HM469003-HM469045, HM469047-HM469064, HM469066-HM469068-HM469077, HM469080, HM469082-HM469088, HM469090-HM469103, HM469105-HM469108, HM469110-HM469144, HM469146-HM469179, HM469181-HM469198, HM469200-HM469213, HM469215-HM469229, HM469231-HM469240, HM469263, HM469271-HM469273, HM469276, HM469278, HM469282, HM469283, HM469285-HM469302, HM469304-HM469306, HM469309, HM469310-HM469312, HM469315, HM469319-HM469321, HM469325, HM469327-HM469329, HM469331-HM469335, HM469337-HM469342, HM469344-HM469353, HM469355, HM469356, HM469359-HM469361, HM469362, HM469364-HM469366, JQ674753-JQ675288. Subtype was determined using the REGA HIV-1 Subtyping Tool 2.0 [21]. Because subtype B predominates in the Canadian epidemic, we limited our analysis to this subtype.

Mixed Base Calls

Mixed bases were identified on sequencing chromatograms at nucleotide positions where a second trace, representing a different base, was present above a threshold τ percentage area of the dominant, primary peak. For example, at a threshold τ = 15%, the secondary peak must represent at least 15% of the primary peak for a mixed base to be called. We varied τ in SeqScape [22] from 5% to 45% for all samples sequenced in our laboratory, creating a new sequence alignment each time, and counted mixed bases with DAMBE [23].

Receiver Operator Characteristic (ROC) Analysis

ROC analysis was used to assess whether sequences could be classified as recent or established on the basis of numbers of mixed bases. We name this method the mixed base classifier (MBC). Under the hypothesis that numbers of mixed bases increase with duration of infection, we first determined with the NHRL training data set the threshold τ that maximized area under the ROC curve (AUC). We then used the full training data set to search for the mixed base cutoff γ155 that maximized sensitivity and specificity in distinguishing recent from long-term infections at 155 days.

Sequence Subsets

The following 11 subsets were created from the full training data set to contain only sites potentially informative for the MBC: (1) sites where numbers of mixed bases increased in established infections (True +); sites where entropy was increased at the (2) nucleotide level (ΔE + Nuc) or (3) amino acid level (ΔE + AA); sites where the entropy increase was significant at the (4) nucleotide level (ΔE + Nuc.Sig) or (5) amino acid level (ΔE + AA.Sig) (Figure 1); (6) sites that were associated with HLA (HLA +); (7) sites that were under positive selection (Pos); (8) sites that were HLA associated and

![Figure 1](image-url). Site-specific differences in entropy (ΔEntropy). Site-specific ΔEntropy values were calculated between recent and established infections in the full training data set, both in nucleotide (A) and in amino acid (B) sequences. Site-specific ΔEntropy values are color-coded: red represents significant differences, and blue represents nonsignificant differences.
under positive selection (HLA + pos); and (9–11) sites at codon positions 1–3. Table 1 lists the resulting alignment sizes (for positions, see Supplementary Data).

Shannon entropy measures the variability at each position. Site-specific differences in entropy between recent and established infections were calculated using the Shannon entropy tool (http://www.hiv.lanl.gov/content/sequence/ENTROPY/entropy.html); significance was assessed by a randomization test (α = 0.05). HLA-associated sites were described previously [24]. Sites under positive selection were inferred with SLAC [25], under the GTR substitution model.

Mixed bases were counted in each subset. ROC analysis, which plots the true-positive rate of the new test (sensitivity) against its true negative rate (calculated as 1 – specificity), was used to identify the mixed base cutoff $\gamma_{155}$ that best separates recent from established infections on the basis of the AUC: the best classifier has the highest AUC. This analysis was performed on each subset, and the best classifier was tested in the BED data set. As viral diversity may depend on transmission route [12], we further compared mixed base frequencies between patients infected through sexual exposure and those infected through blood exposure. All statistical analyses were performed using SPSS [26].

RESULTS

Mixed Base Thresholds \( \tau \) of 15% and 25% Both Perform Well in the MBC

To test whether the MBC could classify infections as <155 days or >155 days old as accurately as it classifies them as <1 year or >1 year old [12], we first assessed the effect of varying the threshold \( \tau \) for calling mixed bases. In the NHRL training data set, we counted mixed bases at mixture thresholds (\( \tau \)) of 5%, 15%, 25%, 35%, and 45% of the dominant peak. The AUC for different thresholds \( \tau \) ranged from 0.702 (95% confidence interval [CI], .597–.806) to 0.802 (95% CI, .706–.897). AUC was highest at thresholds \( \tau = 15\% \) and \( \tau = 25\% \) (Figure 2A), with \( \tau = 15\% \) yielding a sensitivity and specificity of 78.8% and 80%, respectively. At \( \tau = 15\% \), only a small proportion of sequences (14.6%) remained misclassified.

Baseline Performance of MBC on the Full Training Data Set

In the full training data set, chromatograms were not available to vary the threshold \( \tau \), but consistent with our methods, \( \tau \) was known to lie between 15% and 25%. Mixed base frequency distribution differed significantly between recent sequences and established sequences (on average, 0.22% and 1.28%, respectively; \( P < .001 \) by the Kolmogorov-Smirnov [KS] test). ROC analysis indicated that a cutoff $\gamma_{155} = 0.455$ provided the highest sensitivity and specificity (77.8% and 81.9%, respectively), such that sequences containing <0.45% mixed bases should be classified as recent. ROC analysis was repeated after excluding samples from the NHRL training data set, leaving 96 recent and 141 established samples. In this data set, $\gamma_{155} = 0.465$ provided the highest sensitivity and specificity (81.3% and 83%, respectively).

An Entropy-Based Approach Can Increase Sensitivity and Specificity of the MBC

Performance of the MBC was further increased in 4 of the subsets: True +, \( \Delta E + AA \), \( \Delta E + AA.Sig \), and \( \Delta E + Nuc \) (Table 2). Of these, the \( \Delta E + Nuc \) sites, which included only 27.28% of the sequenced nucleotide positions, most improved MBC performance, with sensitivity and specificity increasing to 85.2% and 83.5%, respectively, at a mixed base cutoff $\gamma_{155} = 0.825$. The negative predictive value increased from 0.82 to 0.86, and the positive predictive value increased from 0.80 to 0.83 as compared to use of full sequences. The improvement from using only \( \Delta E + Nuc \) sites appeared highly significant: when 356 sites were randomly sampled without replacement from the full training data set 10 times (among only \( pol \) variable sites), all resulting ROC curves had lower AUCs, with nonoverlapping CIs (Figure 2B).

HLA-Associated Sites and Sites Under Positive Selection Are Not Sufficient to Predict RHI

Because HIV diversifies under selective pressure early in infection [27, 28], we hypothesized that HLA + and Pos sites might increase the predictive power of the MBC. Neither of these categories alone improved the MBC as compared to the full sequence (Table 2), although Pos sites yielded a surprisingly high AUC of 0.802, despite covering only 6.4% of the
alignment. Combination of HLA + and Pos sites increased performance beyond either category alone (AUC = 0.858) but not beyond use of the full sequences. Although both HLA + and Pos sites associated weakly with \( \Delta E+Nuc \) (\( P = .013 \) and \( P = .037 \), respectively, by the Fisher exact test), these sites were insufficient for optimal MBC performance.

**All 3 Codon Positions Are Informative for the MBC**

Codon redundancy results in greater variability in the third codon position; however, restricting analysis to individual codon positions did not improve MBC performance (Figure 3). The best performance was achieved using third codon positions (AUC = 0.872), but there was little difference between codon positions, suggesting that mixed bases resulting from both synonymous and nonsynonymous mutations contribute information for the timing of infection classification using MBC.

**Evaluation of the Concordance Between MBC and BED Results**

Finally, the MBC was compared to BED in the BED data set. Mixed bases in the BED data set were called at a threshold \( \tau = 15\% \). At this threshold, the average proportion of mixed bases was 0.852\% per site for established infections and 0.346\% for recent infections (\( P < .001 \) by the KS test). Using full sequences and \( \gamma_{155} = 0.45\% \), MBC and BED agreed on the classification of 909 of 1354 samples (67.13\%). Of 445 discordant samples, three-fourths were considered established by BED but recent by MBC. When the \( \Delta E+Nuc \) sites determined in the full training data set at \( \gamma_{155} = 0.82\% \) were used, agreement between BED and MBC increased slightly, to 68.98\% (Figure 4). The highest proportion of discordant results remained for specimens classified as established by BED but as recent by MBC.

**No Difference in Mixed Base Frequencies Between Exposure Categories**

Of 1354 specimens in the BED data set, exposure could be categorized as sexual for 368 samples and as blood related for 803 samples. No difference in mixed base frequencies was observed between the 2 groups (0.701\% vs 0.692\%; \( P > .1 \) by the KS test).

**DISCUSSION**

We showed here that, in a precisely timed cohort, the stage of HIV infection can be inferred from the proportion of mixed bases identified during population-based sequencing of the pol region, consistent with previous findings [12, 29–31]. In addition, we improved the MBC and distinguished infections with a duration of <155 days from those with a duration of >155 days, as opposed to the 1-year threshold used by Kouyos et al [12]. This recalibration is crucial because both historical and current estimates of incidence in many Canadian [32] and international studies [33] use the 155-day limit to identify RHI. Because the risk of HIV transmission is elevated early in infection because of high viral loads [2], the ability to classify infections as <155 days old can offer greater value to understanding the dynamics of HIV transmission.

The universal application of MBC depends on calibrating sequencing instruments and base-callers across laboratories [12]. There is no clear standard for the threshold at which mixed bases are called, and both intralaboratory and interlaboratory variability is exacerbated by the manual review of sequences accepted as a component of the genotyping process.
Table 2. Mixed Base Classifier Performance of Each Category of Sites in the Full Training Data Set

<table>
<thead>
<tr>
<th>Alignment name</th>
<th>Full Sequence</th>
<th>True +</th>
<th>ΔE + Nuc</th>
<th>ΔE + Nuc. Sig</th>
<th>ΔE + AA</th>
<th>ΔE + AA. Sig</th>
<th>HLA +</th>
<th>Pos</th>
<th>HLA + Pos +</th>
</tr>
</thead>
<tbody>
<tr>
<td>Best cutoff $\gamma_{155}(%)$</td>
<td>0.45</td>
<td>0.90</td>
<td>0.82</td>
<td>1.00</td>
<td>0.50</td>
<td>0.54</td>
<td>0.71</td>
<td>0.64</td>
<td>1.10</td>
</tr>
<tr>
<td>AUC</td>
<td>0.878</td>
<td>0.899</td>
<td>0.906</td>
<td>0.843</td>
<td>0.890</td>
<td>0.899</td>
<td>0.849</td>
<td>0.802</td>
<td>0.858</td>
</tr>
<tr>
<td>Sensitivity</td>
<td>0.827</td>
<td>0.846</td>
<td>0.852</td>
<td>0.870</td>
<td>0.852</td>
<td>0.889</td>
<td>0.796</td>
<td>0.809</td>
<td>0.870</td>
</tr>
<tr>
<td>Specificity</td>
<td>0.788</td>
<td>0.747</td>
<td>0.835</td>
<td>0.759</td>
<td>0.776</td>
<td>0.782</td>
<td>0.788</td>
<td>0.747</td>
<td>0.712</td>
</tr>
</tbody>
</table>

Four entropy-based measures are listed: sites that increased in entropy in nucleotide (ΔE + Nuc) and amino acid sequences (ΔE + AA), and sites that significantly increased in entropy in nucleotide (ΔE + Nuc.Sig) and amino acid sequences (ΔE + AA.Sig); as well as sites previously shown to be HLA associated (HLA +); sites inferred under positive selection (Pos); and a combination of both (HLA + Pos +).

Abbreviation: AUC, area under the curve.

* Categories of sites that outperformed the full sequence.

Here we showed that mixed base thresholds of 15% and 25%, available in SeqScape [22] and Trugene, perform equally well in the MBC. Because commonly used thresholds lie somewhere between these values, most HIV pol sequences generated should be appropriate for use with MBC, even in the absence of sequencing chromatograms. Moreover, the performance of MBC was not decreased in the full training data set, where the threshold varied between 15% and 25%, as compared to the NHRL training data set, where $\tau$ was fixed for all sequences. Taken together, our results indicate that it may not matter whether the same threshold is used across all sequences analyzed, as long as it falls within this range.

We next examined whether certain sites could offer greater discriminatory power to distinguish recent from established infections. Our analysis revealed that, although the third codon position contained most of the information relevant to the MBC, the predictive power of the MBC was reduced when using individual codon positions. Similarly, although HLA-associated sites are known to diversify rapidly after transmission [27, 28], the exclusive use of these sites did not improve the MBC. It is possible that, although variability at HLA sites initially increases under adaptive immune pressure, continued selective pressure ultimately results in the fixation of better adapted variants [34], reducing numbers of mixed bases at HLA sites. Another possibility is that, because HLA adaptation occurs early in infection [35], mixed bases are already present at HLA sites in RHI, confounding the MBC. However, the performance of MBC improved significantly by focusing on the 25% of sites that increased in entropy (Figure 2B).

Our results add to a growing body of evidence that the MBC may be considered as a potential addition to the public health toolbox for the identification of RHI. A significant advantage of the MBC is that no additional laboratory work is required beyond the genotyping performed for clinical or
surveillance purposes. A freely available MBC has been implemented in DAMBE [23], and the cutoff \( \gamma_{155} \) identified in this study can be directly applied to existing \( pol \) sequences generated for determining subtype and identifying drug resistant mutations. Application of MBC to existing longitudinal sequencing databases may create new potential models of HIV transmission patterns. Indeed, even if our analysis is entirely cross-sectional, we still found significantly higher frequencies of mixed bases among infections with a duration of >155 days \((P < .001)\). One remaining point is the less-than-perfect concordance (70%) between MBC and the BED. The high frequency of mixed bases observed in sequences classified as recent by BED is consistent with the assay’s tendency to overclassify infections as recent [36]. This result may also highlight the fact that high viral diversity is already present in some patients in primary HIV infection, possibly because of the rapid proliferation of viral variants prior to immune selection [37–39]. However, we found the greatest discordance among established infections that had fewer mixed bases. This latter finding may result from the influence of sequences from much-longer-term infections, in which viral diversity decreases. As estimates of BED accuracy vary widely [40, 41], it is possible that the lack of observed concordance on a per-specimen basis reflects the imperfect performance of BED in predicting RHI. Unfortunately, our early infections were serology negative, and determining concordance of the BED and MBC on these specimens is impossible because the BED requires positive serologic findings. This being said, the widespread reporting of BED results in the scientific literature means that reconciling BED results with MBC findings remains a critical, albeit challenging, endeavor. Contrary to Kouyos et al [12], we did not find that viral diversity depended on mode of HIV acquisition.

Another limitation is that the sequences used in this study represent a heterogeneous group of early or late infections. The late infection sequences were obtained from individuals infected anywhere from 156 days to >3 years before sampling. Although all patients in the BED data set had newly diagnosed infection, the stage of infection varied. It is unlikely that any sequences were derived from patients with AIDS because AIDS cases are so infrequent in Canada and mostly occur among patients in whom HIV infection was previously diagnosed [42]. Meanwhile, the early infection group consisted of specimens from patients identified as p24 positive before seroconversion, as well as from individuals classified as having RHI on the basis of discordant results of HIV tests performed up to 155 days apart. This inclusion of sequences from infections prior to seroconversion, when genetic diversity is known to be low, may influence results by favoring the hypothesized association.

We further highlight that the results from this analysis are only applicable to ART-naive sequences, such as those used in this study. Just as patterns of mutation differ in treated patients, suppressed viral loads will make it harder to amplify enough copies to obtain an accurate representation of viral diversity. While samples collected through the SDR exclusively originate from ART-naive patients with a recent diagnosis, equivalent programs in other countries, such as the United Kingdom, continue to collect samples from patients beyond this period.

To fully validate the MBC, it would be necessary to have access to data sets of precisely timed and longitudinal samples, but unfortunately very few such data sets exist to date. Before results can be extrapolated, further research should focus on evaluating the MBC on alternative data sets, such as involving ART-treated subjects, elite controllers, and patients with AIDS, all of which have presented problems for antibody-based incidence assays [4].

In conclusion, we found highly significant differences in mixed base frequencies between samples associated with infection durations of <6 months or >6 months and have identified sites in \( pol \) at which mixed bases most correlate with the time since infection. We argue that the MBC is a significant addition to the tool kit for detecting RHI. In particular, it might be used in combination with other tools, such as the recent infection testing algorithm, which uses results from laboratory assays, as well as clinical information [43]. While future work should validate our list of entropy-identified, highly informative sites and should determine the functional significance of those that were not HLA associated, longitudinal analyses of patients with known HLA types might better determine whether mixed bases do indeed appear at HLA-associated sites. This step will likely be facilitated by the replacement of Sanger sequencing by next-generation sequencing, which is better suited to quantifying population genetic diversity.

**Supplementary Data**

Supplementary materials are available at *The Journal of Infectious Diseases* online (http://jid.oxfordjournals.org/). Supplementary materials consist of data provided by the author that are published to benefit the reader. The posted materials are not copyedited. The contents of all supplementary data are the sole responsibility of the authors. Questions or messages regarding errors should be addressed to the author.

**Notes**

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