Rapid Sequence Change and Geographical Spread of Human Parvovirus B19: Comparison of B19 Virus Evolution in Acute and Persistent Infections

Päivi Norja, Anna Maria Eis-Hübinger, Maria Söderlund-Venermo, Klaus Hedman, and Peter Simmonds

Centre for Infectious Diseases, University of Edinburgh, Summerhall, Edinburgh EH9 1QH, United Kingdom; Institute of Virology, University of Bonn, Sigmund-Freud-Strasse 25, D-53105 Bonn, Germany; and Department of Virology, University of Helsinki, and Helsinki University Hospital Laboratory Division, Haartmaninkatu 3, FI-00290 Helsinki, Finland

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Parvovirus B19 is a common human pathogen maintained by horizontal transmission between acutely infected individuals. However, B19 virus can also be detected in tissues throughout the life of the host, although little is understood about the nature of such persistence. In the current study, we created large VP1/2 sequence data sets of plasma- and tissue (autopsy)-derived variants of B19 virus with known sample dates to compare the rates of sequence change in exogenous virus populations with those in persistently infected individuals. By using linear regression and likelihood-based methods (such as the BEAST program), we found that plasma-derived B19 virus showed a substitution rate of \( 4 \times 10^{-4} \) and an unconstrained (synonymous)-substitution rate of \( 18 \times 10^{-4} \) per site per year, several times higher than previously estimated and within the range of values for mammalian RNA viruses. The underlying high mutation frequency implied by these substitution rates may enable rapid adaptive changes that are more commonly ascribed to RNA virus populations. These revised estimates predict that the last common ancestor for currently circulating genotype 1 variants of B19 virus existed around 1956 to 1959, fitting well with previous analyses of the B19 virus “bioportfolio” that support a complete cessation of genotype 2 infections and their replacement by genotype 1 infections in the 1960s. In contrast, the evolution of B19 virus amplified from tissue samples was best modeled by using estimated dates of primary infection rather than sample dates, consistent with slow or absent sequence change during persistence. Determining what epidemiological or biological factors led to such a complete and geographically extensive population replacement over this short period is central to further understanding the nature of parvovirus evolution.

Parvovirus B19 is the best-characterized member of the family Parvoviridae, known to cause pathogenic infections in humans (14). Parvoviruses are classified into genera or groups with differences in host range, pathology, and tissue/cellular tropism (14). Human parvovirus B19 is a member of the Erythrovirus genus that specifically targets erythroid progenitor cells through the globoside P antigen (2). Infections are associated with anemia and sometimes with aplastic crises.

B19 virus has respiratory, vertical (where infection may lead to intrauterine anemia or fetal death [hydrops fetalis] [4]), and parenteral (often through the contamination of plasma-derived blood products [3]) routes of transmission. Primary infections are characterized by an intense viremia of short duration, a rapid resolution following seroconversion for antibody to B19 virus, and lifelong immunity from reinfection (23). In Western countries, the frequency of B19 virus exposure rises steeply with age in individuals between the ages of 1 year and the late-teenage years to approximately 60 to 70% (13, 28, 33, 44).

Little is known about the origin, past epidemiology, or evolution of B19 virus. However, the lifelong persistence of viral DNA in tissues (11, 15, 19, 22, 30, 41) may reveal what variants circulated in previous decades of life when primary infections occurred (30). There are three genotypes of B19 virus, differing in nucleotide sequence by approximately 13 to 14% (6, 16, 29, 37); genotypes 1 and 2 are found in Europe, the United States, and other Western countries, while genotype 3 is restricted to sub-Saharan Africa and South America (6, 31, 34). Sequences of both genotype 1 and 2 were found in tissues of persistently infected individuals, although genotype 2 detection was strictly limited to those born before 1973, indicating that the circulation of this genotype ceased in Finland and Germany after this date (30). A similar restriction of genotype 2 to those born before 1963 was found in study subjects from Scotland (27), indicating some commonality in the transmission networks and dynamics of B19 over large areas of Europe.

The relatively dynamic picture of B19 virus populations suggested by these findings has reawakened interest in the evolution and molecular epidemiology of B19 virus. To estimate the times of origin of genotype 1 variants that replaced genotype 2 in Europe, we created and analyzed a large data set of genotype 1 VP1/VP2 sequences of plasma-derived \( (n = 33) \) viruses. We additionally studied tissue-derived variants of B19 virus to compare the evolution of horizontally transmitted, exogenous virus populations with that of virus populations occurring dur-
The sequence data set was supplemented with published sequences with annotated sample dates and sources (i.e., plasma or tissue) or provided by the investigators (see Table S3 in the supplemental material).

Screening for recombination and positive selection. All sequences were screened for recombination through the detection of phylogeny violations between trees constructed from sequential 501-base fragments, increasing by 252-base increments through the ORF2 region by using a TreeOrder scan in a Simmonic package. The only violations detected at a bootstrap value of >65% involved the previously described recombinant sequence AN66 (38), which was excluded from further analysis. No evidence for positively selected amino acid residues among the remaining data set of 121 ORF2 sequences was obtained by using CodeML (45) or by using the HyPhy package (all P values were >0.01 [32]).

Phylogenetic analysis and rate calculation. Phylogenetic trees of B19 virus ORF2 sequences were constructed by neighbor-joining of Jukes-Cantor analysis (J-C)-corrected pairwise distances. Regression analysis was performed by using maximum composite likelihood (MCL) distances (calculated by MEGA4 [43] with a gamma distribution value of 0.179) between each sequence and the earliest dated sequence available, PVBPRO. The separate regression analysis of synonymous sites was performed by using J-C-corrected distances.

A Markov chain Monte Carlo (MCMC) method implemented in BEAST package version 1.4 (9) was used to independently estimate the rate of the sequence change of B19 virus (10). Dated sequence sets were run with chain lengths of 10 million under the SRD06 model of substitution (39) and the assumption of a constant population size. Separate runs using a strict molecular clock and a relaxed (uncorrelated lognormal) clock were carried out. All other parameters were optimized during the burn-in period. The output from BEAST was analyzed by using the program TRACER (http://beast.bio.ed.ac.uk/Tracer).

Nucleotide sequence accession numbers. B19 virus sequences were assigned the accession numbers EU478527 to EU478589 (see Table S1 in the supplemental material).

RESULTS

Sequence variability of B19 virus. Sequence diversity within and between genotypes of B19 virus occurs primarily at synonymous sites. We examined further constraints on B19 virus sequence changes other than protein coding in ORF1 and ORF2 by measuring the synonymous-site variability (SSV) within (Fig. 1A) and between (Fig. 1B) genotypes. A region of suppressed SSV was observed in the overlap of ORF1 and the 7-kDa protein translated from a different reading frame (positions 1904 to 2122 [25]) and in the overlap with the 11-kDa protein gene with ORF2 (positions 4704 to 4988 [42, 46]). A third conserved coding sequence in all three genotypes, referred to previously as the “X” ORF (14), was associated with profound suppression of SSV between positions 2650 and 3200. The alternative reading frame was under a greater sequence constraint than ORF2 (Table 1), with lower nonsynonymous-site variability in the “X” ORF than in ORF2 and a lower ratio of nonsynonymous evolutionary changes (dN) to synonymous changes (dS) (<0.3). All regions of ORF2 with

<table>
<thead>
<tr>
<th>Frame</th>
<th>Position</th>
<th>Type of comparison</th>
<th>dS</th>
<th>dN</th>
<th>dN/dS</th>
</tr>
</thead>
<tbody>
<tr>
<td>ORF2</td>
<td>2714–2729</td>
<td>Intrageneotype *</td>
<td>0.0139</td>
<td>0.0096</td>
<td>1.21</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Intergeneotype</td>
<td>0.0733</td>
<td>0.0363</td>
<td>0.59</td>
</tr>
<tr>
<td>“X” ORF</td>
<td>2712–2930</td>
<td>Intergeneotype</td>
<td>0.0261</td>
<td>0.0053</td>
<td>0.28</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Intergeneotype</td>
<td>0.1011</td>
<td>0.0259</td>
<td>0.28</td>
</tr>
</tbody>
</table>

* J-C-Corrected mean pairwise distances.

**Weighted pairwise comparison of sequences of the same genotype.
overlapping coding sequences were excluded from subsequent evolutionary analysis.

**Time-related sequence change of B19 virus ORF sequences.**

Phylogenetic analysis of B19 virus sequences from positions 3242 to 4612 in ORF2 amplified from 33 plasma samples or from archived plasma pools/blood products and 17 published plasma-derived sequences of known collection dates was carried out. This showed evidence for the time-related sequence drift of B19 virus (Fig. 2A), with increasing divergence from the root of the tree with later sampling dates. Regression analysis of nucleotide distances between each sequence and PVBPRO with consideration of the sample collection date (Fig. 3) revealed the rate of sequence change of each sequence data set to be around $4.4 \times 10^{-4}$ substitution per site per year (Table 2). Sequence changes at synonymous sites showed the same time-related correlation ($R^2 = 0.51$) and a much higher substitution rate ($18 \times 10^{-4}$ substitution per site per year). No association between sequence divergence at nonsynonymous sites and time was detected in either data set.

To investigate whether B19 virus genotypes evolved in a
comparable manner during persistence, the same analysis was performed on sequences recovered from tissue samples (Fig. 2). The observable lack of temporal structure in the tree (Fig. 2B) was confirmed by the lack of correlation between the sample date and divergence as determined by regression analysis (4.4 × 10⁻⁴ substitution per site per year) (Table 2), lying within the high-probability distribution (HPD) of the MCMC values. Predicted dates for the most recent common ancestor (MRCA) of the sequence analyzed in the plasma-derived data set were from 1956 to 1959 (the HPD range was 1941/1945 to 1968/1969) (Fig. 2).

The substitution rates calculated for autopsy-derived variants of B19 virus by the MCMC method were 10-fold lower than those calculated for plasma-derived viruses (Table 2). The lower HPD interval was close to zero, consistent with an extremely low consistent rate of or undetectable sequence change among these sequences. These data are consistent with the lack of association between divergence and sample by regression analysis (Fig. 2B).

Rate of sequence change of B19 virus during persistence. The lack of correlation between divergence and sample year observed for tissue-derived B19 virus variants may be accounted for by restricted or absent virus replication during persistence. The immediate problem with investigating this hypothesis is the lack of information about the time of primary infection in the study subjects from whom tissue samples were obtained. However, as a group, their most likely expected times of infection may be inferred from standard incidence and seroprevalence information on B19 virus from the United Kingdom and Finland. Both countries showed a period of peak incidence between 8 and 9 years of age, at which time, overall seroprevalence was 35% (i.e., approximately half that of the adult population) (28).

From the subset of study subjects whose ages at the time of sampling were recorded (see Table S1 in the supplemental material), we compared two models of sequence change. Model A assigns a mean time of infection of 9 years of age and assumes no sequence change during the time that the virus persisted. Model B assumes a sequence change after infection at the same rate as that of the exogenous, horizontally transmitted virus. Model A provides a convincing correlation between the time of infection and the substitution rate at all and at silent sites (Table 3) (R² = 0.32 to 0.33; P ≈ 0.01; substitution rates, 3.2 × 10⁻⁴ and 12.9 × 10⁻⁴ per site per year). Lower correlation coefficients likely arose from a greater scatter of the data associated with the introduction of a further variable (age of infection) and the lower number of subjects analyzed. The average of the measured substitution rates of (3.7 ± 2.0) × 10⁻⁴ and (18.3 ± 8.5) × 10⁻⁴ substitution per site per year at all and at synonymous sites, respectively, were remarkably similar (although with higher standard errors) to estimates based on regression analysis for plasma virus (Table 2). No such association is evident in model B (concordant with the analysis of the larger data set) (Table 2).
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of exogenous and persistent B19 viruses were considered to be equal.

with no subsequent sequence change; for model B, the rates of sequence change

model B, the analysis was restricted to the same subset as that used for model A.

before 1973 were excluded. For study subjects aged 9 years or less, infections

change.

the result of the inclusion of sequences derived from biopsy or

substitution rates calculated.

gene coding VP1/2 (31, 38) substantially reduced the overall

invariant regions in the analysis in previous analyses of the

frame down to close to zero. The inclusion of these relatively

gene sequences which effectively drove SSV in the VP1 coding

regions of suppressed variability in VP1 containing overlapping

(31) and for a short region of VP1 of South American genotype

of 38 genotype 1 sequences (38). The rate is additionally sub-

those calculated for this region by the MCMC method for a set

NA, not applicable. With substitution rates, there was no evidence for time-related sequence change.

a Significant values ($P < 0.05$) are bold.

b Mean rate ± standard error.

c NA, not applicable. There was no evidence for a time-related sequence change.

d Mean value (HPD interval).

**DISCUSSION**

**Substitution rate of B19 virus.** Both the MCMC and regression analyses found consistently high, concordant substitution rates of around $4 \times 10^{-4}$ substitution per site per year for B19 virus in ORF2 (Table 2), approximately fourfold higher than those calculated for this region by the MCMC method for a set of 38 genotype 1 sequences (38). The rate is additionally substantially higher than that calculated for genotype 3 sequences (31) and for a short region of VP1 of South American genotype 1 variants (1).

There are two likely reasons for the observed discrepancies in rates between studies. Previous analyses of ORF2 included regions of suppressed variability in VP1 containing overlapping gene sequences which effectively drove SSV in the VP1 coding frame down to close to zero. The inclusion of these relatively invariant regions in the analysis in previous analyses of the gene coding VP1/2 (31, 38) substantially reduced the overall substitution rates calculated.

The second reason for lower estimates in previous studies is the result of the inclusion of sequences derived from biopsy or autopsy tissue samples (e.g., 11 of 38 VP1/2 sequences analyzed by Shackleton and Holmes [38]). However, we have shown that sequence change in B19 virus variants during the time of virus persistence is not modeled adequately by assuming that they have the same substitution rate as horizontally transmitted viruses (Table 3, model B). To demonstrate the effect of their inclusion, we analyzed a combined data set of plasma- and tissue (autopsy)-derived B19 virus variants by the MCMC method and obtained a composite substitution rate for B19 virus of $1.66 \times 10^{-4}$ (HPD, $1.17 \times 10^{-4}$ to $2.19 \times 10^{-4}$) substitution per site per year, three times lower than the rate based on plasma-derived viruses. This figure would be even lower if regions of suppressed variability in VP1 had been included in the analyzed sequences.

**Comparison with substitution rates of other viruses.** The sequence change in VP1/2 and elsewhere in the B19 virus genome is constrained by extreme conservatism in its amino acid sequence of encoded viral proteins (14). Almost all sequence change (even between the more divergent genotypes of B19 virus) occurs at synonymous sites, with extremely low $dN/dS$ ratios for both major genes (31, 38). A fairer comparison of the underlying frequency and fixation of mutations in the B19 virus population with those of other viruses (whose protein sequence may not be under such constraint or may indeed be subject to positive selection) may be achieved by comparing synonymous-substitution rates. In the current study, regression analysis was allowed to calculate an unconstrained-substitution rate for B19 virus of at least $1.8 \times 10^{-3}$ synonymous substitution per site per year (Table 2). This rate lies within the range (of substitutions per site per year) reported for flaviviruses ($0.4 \times 10^{-3}$ to $4.9 \times 10^{-3}$ [restricted to nonzero rates]), paramyxoviruses ($0.7 \times 10^{-3}$ to $4.8 \times 10^{-3}$), and orthomyxoviruses ($0.4 \times 10^{-3}$ to $4.2 \times 10^{-3}$) but was higher than ranges reported for togaviruses and reoviruses ($0.1 \times 10^{-3}$ to $0.78 \times 10^{-3}$ and $0.6 \times 10^{-3}$ to $1.4 \times 10^{-3}$, respectively) and lower only than those reported for picornaviruses ($2.5 \times 10^{-3}$ to $12 \times 10^{-3}$).

The factors that contribute to the rapid sequence drift of

**TABLE 2. Estimates of the rates of sequence change of B19 viruses by using regression and BEAST analyses**

<table>
<thead>
<tr>
<th>Source of B19</th>
<th>Origin of sequence set</th>
<th>Regression analysis (MCL distance)</th>
<th>MCMC analysis (BEAST)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Total no. of samples</td>
<td>$R^2$</td>
</tr>
<tr>
<td>Plasma</td>
<td>This study</td>
<td>33</td>
<td>0.497</td>
</tr>
<tr>
<td></td>
<td>Published sequences</td>
<td>17</td>
<td>0.442</td>
</tr>
<tr>
<td>Combined</td>
<td>All</td>
<td>50</td>
<td>0.547</td>
</tr>
<tr>
<td>Autopsy tissue</td>
<td>This study</td>
<td>29</td>
<td>0.039</td>
</tr>
<tr>
<td></td>
<td>Published</td>
<td>17</td>
<td>0.300</td>
</tr>
<tr>
<td>Combined</td>
<td>Combined Synonymous</td>
<td>46</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td>Combined Nonsynonymous</td>
<td>46</td>
<td>0.014</td>
</tr>
</tbody>
</table>

$^a$ Significant values ($P < 0.05$) are bold.

$^b$ Mean rate ± standard error.

$^c$ NA, not applicable. There was no evidence for a time-related sequence change.

$^d$ Mean value (HPD interval).

**TABLE 3. Comparison of two models of B19 sequence change during persistence**

<table>
<thead>
<tr>
<th>Model$^a$</th>
<th>Total no. of study subjects$^b$</th>
<th>All sites</th>
<th>Synonymous sites</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$R^2$</td>
<td>Rate ($10^{-4}$)</td>
<td>$R^2$</td>
</tr>
<tr>
<td>A</td>
<td>0.330</td>
<td>3.2 ± 1.1</td>
<td>0.318</td>
</tr>
<tr>
<td>B</td>
<td>0.000</td>
<td>0.95</td>
<td>NA</td>
</tr>
</tbody>
</table>

$^a$ For model A, infection was considered to have occurred at the age of 9 years, with no subsequent sequence change; for model B, the rates of sequence change of exogenous and persistent B19 viruses were considered to be equal.

$^b$ For model A, Excludes study subjects with estimated dates of infection before 1973 were excluded. For study subjects aged 9 years or less, infections assumed to have occurred halfway between birth and the sampling time. For model B, the analysis was restricted to the same subset as that used for model A.

$^c$ NA, not applicable. There was no evidence for a time-related sequence change.

$^d$ Boldface values are significantly different.

The factors that contribute to the rapid sequence drift of
B19 virus and potentially to the drift of other parvoviruses maintained by long, horizontal transmission chains are unknown. B19 virus, in common with other small viruses, is replicated by host cell DNA polymerase, and the enzyme’s intrinsic proofreading activity might be expected to reduce the frequency of nucleotide misincorporations to that of host DNA. B19 virus, in common with other small viruses, is replicated by host cell DNA polymerase, and the enzyme’s intrinsic proofreading activity might be expected to reduce the frequency of nucleotide misincorporations. In primates, this mutation frequency leads to a synonymous-substitution rate of approximately $2 \times 10^{-9}$ substitution per site per year (5), around a million times slower than that calculated for B19 virus. The difference in substitution rates cannot be entirely attributed to differences in the number of replicative cycles between B19 virus and its human host. Human DNA in the germ line is copied approximately 30 (females) and 200 (males) times per generation, an average of around 5 times per year. B19 virus cannot possibly undergo 5 million sequential genome replications in a year. With a minimum replication cycle in cells of at least 24 h, during which time at most 10 to 20 sequentially templated copies of the viral genome during the synthesis of concatamerized replication intermediates are likely to occur (8), there would be a maximum of 7,000 sequential genome transcriptions per year. This total is nearly 1,000 times fewer than that expected if B19 virus and human DNA replication occurred with the same underlying mutation frequency.

It is possible that the unidirectional synthesis of DNA during the rolling hairpin replication of parvovirus DNA, resolution, and synthesis of single-stranded genomic DNA sequences for packaging (8) may be associated with higher error frequencies than the bidirectional synthesis of host genomic DNA. Interestingly, substitution rates in RNA viruses are correlated with genome size, with lower rates occurring in viruses with larger genomes (18). This provides evidence that viruses, in common with larger organisms, have the ability to modulate mutation rates to provide the correct balance between replicative fitness and adaptability. One might speculate that some small DNA viruses, such as B19 virus, may have a similar ability to modify the mutation rates of the host cell, perhaps because its transmission and epidemiology, and thus its evolutionary constraints and pressures, might closely resemble those of respiratory viruses with RNA genomes. How this might be achieved is not known.

**Recent spread of parvovirus B19 virus.** Revised estimates for the substitution rates of B19 virus predict a relatively recent date for the MRCA of genotype 1 (Europe, the United States, Japan, Korea, and Brazil). An MRCA occurring in the late 1950s is consistent with the epidemiology of genotype 1 and genotype 2. The large-scale surveillance of B19 virus genotypes in Europe showed that genotype 1 infections are almost invariably detected in contemporary samples, with extremely few if any genotype 2 (or 3) infections (6, 7, 21). This contrasts with the frequent detection of genotype 2 variants in tissue samples or persistently infected (immunosuppressed) individuals (16, 24, 27, 30, 36).

The hypothesis that these variants originate from primary infections when genotype 2 was the predominant circulating virus (30) is supported by the unusual age restrictions of type 2-infected subjects (those born before 1973 and 1963 [27, 30]). These observations are fully consistent with the predicted MRCA of genotype 1 in the late 1950s, which then almost entirely replaced genotype 2 by the early 1970s over large parts of the world. The epidemiological factors and the possible role of biological differences between genotypes, such as greater infectivity or transmissibility, that may have contributed to its dramatic spread and replacement of other genotypes are entirely unknown (12).

**Nature of B19 virus sequence change during persistence.** The observation that the sequence divergence of tissue-associated variants showed no correlation with sample collection dates demonstrates directly that the evolution of B19 virus during its persistence differs from that of horizontally transmitted viruses. However, paired samples from acute infection and after a period of persistence were not available to directly investigate the sequence change, and this study had to rely on estimates of times of infection based on those determined by previous seroepidemiological studies (13, 28, 33, 44). However, the crude model used in the current study (primary infections at the age of 9 and no subsequent sequence change during persistence) actually fitted the evolutionary data closely and restored the correlation between sequence divergence and the estimated year of primary infection (Table 3).

Models A and B compared in this study are extremes, and the observed better fit of model A to the data clearly does not rule out other scenarios, such as the persistence of the virus with a much lower rate of sequence change arising from restricted virus replication or ongoing rapid sequence change in a small subset of individuals and an absence in the majority. Observations for maintained strong cytotoxic-T-cell reactivity in B19 virus-seropositive individuals (17) and low-level fluctuating viremia in 0.3 to 0.8% of healthy blood donors (20, 35) do indeed imply various degrees of ongoing replication in vivo. It remains uncertain whether individuals with reactivated virus may be infectious; it is therefore possible that the infrequent cases of genotype 2 infections recorded in Europe (7, 21, 24) may have arisen by the transmission of reactivated virus from persistently infected individuals rather than by the continued low-level circulation of this genotype in the community. Understanding the nature of the persistence of B19 virus is therefore important for future investigations of its molecular epidemiology and evolution and of the factors underlying the large-scale population replacements that we have witnessed in the past few decades.

**ACKNOWLEDGMENTS**

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