- Selective whole genome amplification as a tool to enrich specimens with low *Treponema pallidum* genomic DNA copies for whole genome
 sequencing
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16 Running title: Sequencing of *T. pallidum* from Clinical Specimens

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19

20 Abstract.

21 Downstream next generation sequencing of the syphilis spirochete Treponema pallidum subspecies pallidum (T. pallidum) is hindered by 22 low bacterial loads and the overwhelming presence of background metagenomic DNA in clinical specimens. In this study, we investigated selective whole genome amplification (SWGA) utilizing Multiple Displacement Amplification (MDA) in conjunction with custom 23 oligonucleotides with an increased specificity for the T. pallidum genome, and the capture and removal of CpG-methylated host DNA followed by 24 25 MDA as enrichment methods to improve the yields of T. pallidum DNA in rabbit propagated isolates and lesion specimens from patients with 26 primary and secondary syphilis. Sequencing was performed using the Illumina MiSeq v2 500 cycle or NovaSeq 6000 SP platform. These two 27 enrichment methods led to 93-98% genome coverage at 5 reads/site in 5 clinical specimens from the United States and rabbit propagated isolates, 28 containing >14 T. pallidum genomic copies/ μ l input for SWGA and >129 genomic copies/ μ l for CpG methylation capture with MDA. Variant 29 analysis using sequencing data derived from SWGA-enriched specimens, showed that all 5 clinical strains had the A2058G mutation associated 30 with azithromycin resistance. SWGA is a robust method that allows direct whole genome sequencing (WGS) of specimens containing very low 31 numbers of *T. pallidum*, which have been challenging until now.

32 **Importance**

33	Syphilis is a sexually transmitted, disseminated acute and chronic infection caused by the bacterial pathogen Treponema pallidum
34	subspecies pallidum. Primary syphilis typically presents as single or multiple mucocutaneous lesions, and if left untreated, can progress through
35	multiple stages with varied clinical manifestations. Molecular studies rely on direct amplification of DNA sequences from clinical specimens;
36	however, this can be impacted by inadequate samples due to disease progression or timing of patients seeking clinical care. While genotyping has
37	provided important data on circulating strains over the past two decades, whole genome sequencing data is needed to better understand strain
38	diversity, perform evolutionary tracing, and monitor antimicrobial resistance markers. The significance of our research is the development of a
39	SWGA DNA enrichment method that expands the range of clinical specimens that can be directly sequenced to include samples with low numbers
40	of T. pallidum.
41	Introduction
42	Syphilis rates have been steadily increasing in the United States with 38,992 cases (11.9 per 100,000 people) of primary and secondary
43	syphilis and 1,870 cases (48.5 per 100,000 live births) of congenital syphilis reported to the CDC during 2019 (1). This represents a 167.2%
44	increase in primary and secondary syphilis rates since 2010 and a 291.1% increase in congenital syphilis reported since 2015. While syphilis rates
45	have been on the rise in the U.S., the genetic diversity of the bacterial pathogen Treponema pallidum subspecies pallidum (hereafter referred to as
46	T. pallidum), in this setting, is not well understood due to the lack of recently sequenced whole genomes from clinical specimens. Strain diversity
47	has been gleaned from molecular epidemiology studies, which are based on 3 to 4 genetic loci, but may not be representative of the entire T.
48	pallidum genome (2-5).
49	Molecular studies have relied primarily on T. pallidum strains propagated in rabbits or DNA amplified directly from clinical specimens,

50 because *T. pallidum* cannot be grown on routine laboratory media. However, advances have been made with *in vitro* tissue culture and the

51 propagation of *T. pallidum* in rabbits from cryopreserved genital lesion specimens, which may make routine culture directly from clinical 52 specimens a possibility in the near future (6-7). Despite these advances, the methods are still time-consuming and impractical for laboratory

53 diagnosis and molecular epidemiological studies of syphilis.

54 Metagenomic shotgun sequencing approaches have made significant advances in recent years with sequence data being used for pathogen detection, in silico or whole genome typing, and antimicrobial resistance marker detection, in addition to phylogenetic analyses (8-10). However, 55 56 direct whole genome sequencing (WGS) of T. pallidum from clinical specimens and rabbit isolates can be problematic due to bacterial genomic 57 DNA being outweighed by either human or rabbit DNA. Several DNA enrichment methods have been described for T. pallidum including RNA bait capture techniques, methyl-directed enrichment using the restriction nuclease DpnI, and pooled whole genome amplification, which have 58 59 generated T. pallidum specific WGS data from over 700 metagenomic samples; however, specimens with low numbers of T. pallidum remains challenging (11-16). Therefore, additional approaches that would enable sequencing of samples with low bacterial loads are needed. 60 Azithromycin has been used as an alternative to penicillin for treating early syphilis in the US; however, macrolide-resistant T. pallidum 61 strains, associated with two mutations (A2508G, A2509G) in the 23S rRNA genes, have been reported in many states (17-18). While macrolides 62 are no longer recommended for treatment of syphilis in the US, periodic monitoring is useful to determine the prevalence of resistant strains (19). 63 64 In this study, we describe a robust DNA enrichment method based on selective whole genome amplification (SWGA) using multiple 65 displacement amplification (MDA) and custom primers that enables WGS of clinical specimens with very low genomic copies of T. pallidum and use of the sequence data for macrolide mutation analysis. We also investigated an alternative method that uses CpG methylated capture of host 66 DNA followed by MDA with random oligonucleotide primers. 67

68 **Results**

Real-time qPCR on clinical specimens and spiked samples. The T. pallidum PCR results for all clinical specimens are shown in Table 1. Out of 69 the 11 Emory specimens processed using the standard extraction protocol, only one specimen exceeded 100 genomic copies/µl based on polA 70 qPCR (Table 1). The remaining 10 specimens had an average copy number <1 copy/µl of DNA extract. These 11 specimens had an average 71 72 standardized RNP cycle threshold (RNP_{ct}) value of 30.71 ± 0.13 , and the lowest C_t value (highest concentration of RNP) was 25.22. Based on this data, an RNP_{Ct} value of 25.22 was targeted as the cut-off for the spiked samples below. 73 74 **NEBNext microbiome enrichment with MDA.** The serially diluted spiked samples enriched with the NEB Microbiome Enrichment Kit with 75 subsequent REPLIg Single Cell MDA (hereafter referred to as NEB+MDA) showed a marked increase in *polA* copy number by qPCR (Fig. 1, Table S1). The non-diluted samples indicated an average *polA* copy number of 6.67 x $10^6 \pm 2.74 \times 10^5$ per µl of enriched DNA, which was 603.02 76 77 times greater than the input copy number. The 10-fold diluted samples also indicated increases in *polA* copy numbers, with an average of 7.85 x $10^5 \pm 3.79 \times 10^4$, $1.28 \times 10^5 \pm 1.27 \times 10^4$, $8.66 \times 10^3 \pm 2.54 \times 10^3$, and 964 ± 574.23 copies/µl from 1:10 -1:10,000 dilution, respectively (Table S1). 78 This was a 482 - 995.09 times enrichment when compared to the input copy number. Upon comparing the average RNP_{ct} of each dilution in the 79 80 series, the enriched samples indicated 29.28 ± 1.07 , 31.15 ± 0.46 , 30.25 ± 0.56 , 31.08 ± 0.59 , 31.42 ± 0.45 for the neat -1:10,000 dilution, respectively (Table S1). The RNP_{ct} value of each enriched sample in the dilution series were insignificantly different from one another, with an 81 average $RNP_{Ct} = 30.64 \pm 0.33$ for all dilutions in the series (P = 0.22). 82 83 After enriching with NEB+MDA, the average DNA percent for the neat to 1:10,000 dilutions indicated a range of $2.33\% \pm 0.10$ - 3.91 x

84 $10^{-4}\% \pm 2.50 \text{ x } 10^{-4}\%$ of the total DNA belonging to *T. pallidum*, respectively (Fig. 2). Further, this form of enrichment generated up to a 26.12-

fold increase in the percent of *T. pallidum* DNA, and an average of 16.27-fold \pm 1.92-fold increase, amongst all enriched replicates when compared to the unenriched input. All samples enriched by NEB+MDA were significantly different in their percent *T. pallidum* DNA when compared to their respective inputs (P < 0.01). Apart from enriched samples from the 1:100 and 1:1,000 diluted *polA* inputs, we observed that by increasing the *polA* input copy number 10-fold resulted in a significant increase in the total DNA belonging to *T. pallidum* post-enrichment (P =

89 0.06 and P < 0.05, respectively).

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90 Genome sequencing data derived from samples enriched by NEB+MDA showed 0.01 - 10.52% of the quality-controlled reads binned as 91 T. pallidum, along with a mean mapping read depth to T. pallidum Nichols reference genome ($NC_021490.2$) ranging from 0.05 – 501.75. An average percent coverage of 99.99%, 99.99%, and 97.29% across the Nichols reference genome with at least 5 reads mapped per site (5X) for the 92 93 neat, 1:10, and 1:100 diluted samples, respectively, was observed among the NEB+MDA enriched samples (Fig.3A; Table S1 and Fig. S1). The coverage estimates indicated low deviations from this average in all replicates, with $2.92 \times 10^{-4} \% - 1.38\%$ standard error between all replicates for 94 the neat -1:100 diluted samples. At the same time, for a higher coverage of at least 10 reads mapped per nucleotide (10X), the 1:100 diluted 95 96 samples had an average percentage coverage of 84.14% while neat and 1:10 dilution samples were covered at 99.99% and 99.99% across the 97 reference genome, respectively. A sharp decline in coverage was observed in the 1:1,000 diluted samples, with a break down in replication at an 98 average coverage of 27.08% \pm 18.62% for the 1:1,000 dilution and 4.80% \pm 0.63% for the 1:10,000 diluted samples at 5X read depth. With the QC criteria for efficiency set at \ge 90% at \ge 5X read depth, samples sequenced post NEB+MDA enrichment had a limit of detection (LoD) of 129 99 polA copies/µl of extract (Fig. 3A; Table S1 and Fig. S1). 100

101 Post NEB+MDA enrichment of isolate CDC-SF003, we observed $2.39 \times 10^6 \pm 1.35 \times 10^5$ *polA* copies/µl of DNA extract. Further, 1.06% 102 of the total DNA belonged to *T. pallidum* post enrichment and 3.29% of the host removed quality-controlled sequencing reads were classified as *T*. bioRxiv preprint doi: https://doi.org/10.1101/2021.07.09.451864; this version posted July 10, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. This article is a US Government work. It is not subject to copyright under 17 USC 105 and is also made available for use under a CC0 license.

pallidum. Sequencing indicated a 98.60% coverage across the *T. pallidum* SS14 reference genome (*NC_021508.1*) at 5X read depth with a mean
 mapping depth of 46.43 (Fig. 4; Table 2 and Fig. S2).

105	SWGA Enrichment of <i>T. pallidum</i> Nichols. A total of 12 primer sets were tested by SWGA using Equiphi29 MDA (Table S2-S3). The 1:100
106	diluted Nichols DNA sample (~129 copies/µl) was used to evaluate each of the 12 primers since it was comparable to specimen EUHM-004,
107	which had 106.7 <i>polA</i> copy/ μ l (Table 1; Table S4). Each of the primer sets indicated a 6.86 – 1.16 x 10 ⁵ times enrichment when compared to the
108	input Nichols copy number (Fig. 5A). Further, we observed a >10,000-fold increase in <i>polA</i> copy number in samples enriched with 7 of the 12
109	primer sets (SWGA Pal 2, 4, 5, 9, 10, 11, and 12). SWGA Pal 9 and Pal 11 gave the highest enrichment at 1.13 x 10 ⁵ , and 1.16 x 10 ⁵ times,
110	respectively (Table S4). The difference observed between Pal 9 and Pal 11 in the T. pallidum polA copy number and relative percent DNA
111	belonging to <i>T. pallidum</i> was insignificant; however, Pal 11 was selected for testing the SWGA limit of detection $(P > 0.1; Fig. 5 and Table S4)$.
112	To determine the SWGA Pal 11 primer set's LoD and enrichment for T. pallidum, SWGA was performed in triplicate on the 10-fold
113	dilution series. The $\sim 1.11 \times 10^4$ copies/µl (neat) sample was eliminated from the dilution series, as this was ~ 100 -fold increase in <i>T. pallidum</i> copy
114	number when compared to the clinical specimens tested. We observed a marked increase in polA copy number in every dilution in the series post
115	enrichment (Fig. 1; Table S1). The <i>polA</i> copy number ranged from $1.11 \times 10^6 \pm 6.68 \times 10^5$ for the 1:10,000 dilution to 2.04 x $10^7 \pm 1.20 \times 10^7$ in
116	the 1:10 dilution (Table S1). When compared to the input <i>polA</i> copy number, this was a 2.01 x 10 ⁴ -fold, 1.19 x 10 ⁵ -fold, 3.53 x 10 ⁵ -fold, and 5.53
117	x 10 ⁵ -fold increase in the enriched samples, from 1:10 -1:10,000 dilution, respectively. Upon comparing the average RNP _{Ct} of each dilution in the
118	series, the SWGA enriched samples indicated a $29.36 \pm 0.37 - 28.65 \pm 0.16$ for the 1:10 -1:10,000 dilution, respectively (Table S1). The average
119	RNP_{Ct} at each 10-fold increase in <i>polA</i> concentration were insignificantly different from one another (P > 0.1); however, by increasing the <i>polA</i>
120	input 100-fold, we observed a significant decrease in RNP concentration ($P < 0.03$).

121	After enriching with SWGA, we observed that dilutions ranging from 1:10 to 1:10,000 held 27.93% \pm 1.57% - 3.29% \pm 1.93% of the total
122	DNA belonging to <i>T. pallidum</i> , respectively (Fig. 2). This reflected up to a 1.63 x 10 ⁵ -fold increase in the relative <i>T. pallidum</i> and an average of
123	2.43×10^4 -fold $\pm 1.05 \times 10^4$ -fold increase amongst all replicate SWGA enriched samples when compared to the unenriched samples. All samples
124	were significantly increased in their relative <i>T. pallidum</i> DNA when compared to their respective inputs ($P < 0.0001$). While there was observed
125	deviations in the percent DNA between replicates, the 1:10,000 diluted replicates still yielded a 28.40-fold \pm 17.71-fold increase in DNA
126	belonging to <i>T. pallidum</i> post SWGA when compared to the non-enriched neat dilution ($P < 0.0001$).
127	Genome sequencing data derived from the SWGA enriched Nichols samples showed 0.98%-78.05% of the quality-controlled reads binned
128	as <i>T. pallidum</i> , along with a mean mapping read depth to <i>T. pallidum</i> Nichols reference genome ranging from $65.82 - 4.89 \times 10^3$. An average
129	percent coverage of 98.67% \pm 0.005%, 98.62% \pm 0.003%, and 96.15% \pm 0.082% across the Nichols genome at 5X read depth was observed
130	among the SWGA enriched 10-fold dilution series samples for the 1:10, 1:100 and 1:1,000 diluted samples, respectively (Fig. 3B; Table S1 and
131	Fig. S3). Further, coverage indicated low deviations from this average in all replicates, with a 0.0002% - 1.72% standard error between all
132	replicates for the $1:10 - 1:1,000$ diluted samples. We did observe a sharp decline in coverage from the $1:1,000$ to $1:10,000$ dilution with an average
133	coverage of $38.46\% \pm 2.50\%$ for the 1:10,000 diluted replicates a 5X read depth (Fig. 3B; Table S1 and Fig. S3).
134	Upon comparing the percent T. pallidum DNA derived from both enrichment methods, we observed that SWGA consistently produced
135	higher relative T. pallidum DNA in all samples (Fig. 2). We observed that the 10-fold dilutions enriched with SWGA exhibited an average of
136	94.08-fold - 1.41 x 10 ⁴ -fold increase in relative <i>T. pallidum</i> DNA in the 1:10-1:10,000 diluted samples when compared to the dilutions enriched by
137	NEB+MDA. All dilutions of each enrichment were significantly different from one another ($P < 0.01$), apart from the 1:10,000 and 1:1,000 diluted
138	samples enriched by SWGA and the neat diluted samples enriched by NEB+MDA ($P > 0.07$).

139	Comparing the sequencing data derived from the 1:10 and 1:100 diluted Nichols samples enriched using the NEB+MDA and SWGA, all
140	samples exhibited >95% coverage at 5X read depth (Fig. 3; Table S1 and Fig. S1, S3). There was a decline in coverage observed in the 1:1,000
141	diluted samples enriched by NEB+MDA, with an average coverage of 27.08% \pm 24.80% at 5X read depth. This drop was not observed in the
142	1:1,000 diluted samples enriched by SWGA, which still held >95% coverage at 5X read depth. The 1:10,000 diluted samples enriched NEB+MDA
143	and SWGA exhibited <95% coverage at 5X read depth.
144	Enrichment of Clinical Strains.
145	Due to the increased sequencing coverage derived from the SWGA enriched Nichols strain, SWGA was chosen for enriching clinical
146	specimens with low numbers of <i>T. pallidum</i> (Fig. 3, Table S1). SWGA on clinical specimen EUHM-004 gave an average <i>polA</i> of $6.37 \times 10^6 \pm 2.24$
147	x 10 ⁵ copies/µl with 5.56% of the total DNA belonging to <i>T. pallidum</i> (Table 2). Next generation sequencing using the MiSeq v2 (500 cycle)
148	platform revealed 95.13% coverage across the T. pallidum genome at 5X read depth (Fig. 4; Table 2 and Fig. S2). After large-scale DNA
149	extraction, we observed 31.5 ± 0.5 , 122 ± 1.15 , and 103 ± 6.55 polA copies/µl for specimens EUHM-012 – EUHM-014, respectively (Table 1).
150	For specimen EUHM-012, we observed an average <i>polA</i> of 2.14 x $10^6 \pm 2.82 \times 10^4$ copies/µl with 1.72% of the total DNA belonging to <i>T</i> .
151	pallidum post-enrichment by SWGA (Table 2). Sequencing indicated a 93.98% coverage across the T. pallidum genome at 5X read depth (Fig. 4;
152	Table 2 and Fig. S2).
150	When compared to EUHM 012, EUHM 012 had a higher not a convergence of 5.16 x $10^6 \pm 2.20$ x 10^5 conjectul with 15.480/ of the total

When compared to EUHM-012, EUHM-013 had a higher *polA* copy number at 5.16 x 10⁶ \pm 2.20 x 10⁵ copies/µl with 15.48% of the total DNA belonging to *T. pallidum* (Table 2). The sequencing data correlated with the qPCR data, indicating a 98.56% coverage across the *T. pallidum* genome at 5X read depth (Fig. 4; Table 2 and Fig. S2). We also observed EUHM-014 held an increased *polA* copy number post-SWGA, with 2.57 x 10⁶ \pm 2.21 x 10⁵ copies/µl and 4.72% of the total DNA belonging to *T. pallidum* (Table 2). Upon sequencing, we observed 98.49% coverage bioRxiv preprint doi: https://doi.org/10.1101/2021.07.09.451864; this version posted July 10, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. This article is a US Government work. It is not subject to copyright under 17 USC 105 and is also made available for use under a CC0 license.

157	across the <i>T. pallidum</i> genome at 5X read depth (Fig. 4; Table 2 and Fig. S2). The <i>polA</i> copy number for specimen STLC-001 was 7.42 x $10^6 \pm$
158	7.20 x 10 ⁵ copies/µl with 8.34% of the total DNA belonging to <i>T. pallidum</i> (Table 2). The sequencing coverage was 95.94% at 5X read depth
159	where 38.91% of the quality-controlled reads binned as <i>T. pallidum</i> along with a mean depth read coverage of 1,133.43X (Fig. 4; Table 2 and Fig.
160	S2).

161 Phylogenetic Analysis and Characterization of Genotypic Macrolide Resistance.

162 To analyze whether genomes generated from the 7 clinical specimens or isolates clustered to any of the two deep-branching monophyletic T. pallidum lineages, Nichols-like and Street-14(SS14)-like, a whole genome phylogenetic tree was constructed using the genomes derived from 163 the clinical specimens/isolates along with 126 high quality published T. pallidum genome sequences as of May 2021 (12-15, 20-22; see Table S5, 164 methods in supplemental materials). Phylogenetic analysis revealed the presence of two dominant lineages, of which most strains belonged to the 165 166 SS14-like lineage. We identified a total of four monophyletic clades within this phylogenetic tree with > 30 bootstrap support (Fig. 6). Three of the clinically derived genomes from Atlanta, EUHM-004 (2019) EUHM-012 (2019), and EUHM-014 (2020), belonged to Nichols-like lineage (clade 167 1; n=12; Fig. 6). Interestingly, the other nine Nichols-like genomes in clade 1 were recent clinically derived genomes from Cuba (n=2; 2015-168 169 2016), Australia (n=1; 2014), France (n=2; 2012-2013) and UK (n=3; 2016), and were distinct from the original Nichols strain isolated in 1912 170 and sent to different North American labs as in vivo derived clones, suggesting that we might not yet fully understand the current diversity of this 171 lineage. The three clinical specimens from Atlanta (EUHM-004, EUHM-012, and EUHM-014) and three clinically derived genomes from UK 172 isolated in 2016 (NL14, NL19 and NL17) carried the 23S rRNA A2058G mutation that confers macrolide resistance, suggesting a recent 173 acquisition of this antibiotic resistance variant in the Nichols-like lineage.

174	Even though previous phylogenomic analyses indicated that SS14-lineage showed a polyphyletic structure, our phylogenetic analysis with
175	a greater number of genomes showed the presence of 3 monophyletic clades (Clades 2, 3 and 4)(12, 14; Fig. 6). Clades 2 and 4 contained genomes
176	clustered within the previously reported SS14 Ω -A sub cluster, which also contained two clades corresponding to the clades 2 and 4 detected in this
177	study, and contained genomes derived from Europe and North America; while clade 3 was similar to sub cluster SS14Ω-B and composed of
178	Chinese and North American derived T. pallidum genomes. The rabbit-derived clinical isolate, CDC-SF003 (San Francisco, U.S; 2017) sequenced
179	in this study, clustered within clade 2; while EUHM-013 (Atlanta, U.S; 2020) and STLC-001 (St. Louis, U.S; 2020) genomes clustered within
180	clade 4. Sequence analysis showed that all 3 strains carried the A2058G AMR variant for macrolide resistance. Macrolide resistance strains were
181	widespread among the SS14-lineage with higher proportion among the genomes in clades 2 and 3 compared to clade 4 genomes. The A2058G
182	point mutation identified in 4 patient specimens and isolate CDC-SF003 was verified by real-time PCR testing of genomic DNA and SWGA-
183	enriched samples (data not shown). There was inadequate sample for the fifth specimen to confirm the mutation by real-time PCR testing.
184	All the Nichols-like genomes derived from the NEB+MDA and SWGA 10-fold dilution series that contained T. pallidum reads mapped to
185	\geq 90% of the genome with at least 5X read depth formed a tight monophyletic clade (bootstrap support of 88/100) and clustered with the lab-
186	derived Nichols-Houston-J genome (bootstrap support of 100/100), indicating that genomes generated from both methods are adequate to capture
187	genetic variants required to perform a high resolution phylogenetic analysis (Fig. S4).

188 **Discussion**

WGS of *T. pallidum* is often challenging due to low bacterial loads or the difficulty of obtaining adequate samples for testing. In this
 study, we sought to develop a method for performing WGS from rabbit propagated isolates and clinical specimens containing lower *T. pallidum* numbers, leading us to investigate CpG capture and SWGA.

192 CpG capture has been successfully used for enriching bacterial genomic DNA in metagenomic samples (23-24), but this method has not 193 been used for T. pallidum. During our testing, we observed increases in polA copy numbers and relative T. pallidum percent DNA in the neat to 194 1:1,000 dilutions enriched by NEB+MDA when compared to the non-enriched inputs. Further, the results of the percent T. pallidum observed in 195 the enriched 1:10,000 diluted samples correlated with the decrease in overall coverage across the Nichols genome. Even though we observed an increase in both *polA* copy number and relative percent T. *pallidum* DNA for the enriched diluted 1:1,000 diluted samples, we still only gained 196 197 ~50% genomic coverage. This could be due to the remnant human DNA that was not initially captured prior to MDA, or the loss of T. pallidum DNA during the enrichment. While there was no significant difference in the relative human RNP copy number from dilution to dilution, there is a 198 minimum T. pallidum copy number input required to outweigh the remnant human DNA during the metagenomic shotgun sequencing. Taking the 199 above into consideration, we observed that >129 polA copies/ μ l can generate >95% coverage at 5X read depth from the Nichols strain post 200 201 NEB+MDA. The results observed post NEB+MDA enrichment of clinical isolate CDC-SF003 correlated with the Nichols limit of detection 202 validation, with >98% coverage at 5X read depth across the T. pallidum genome. In silico variant analysis correlated with real-time PCR detection 203 of the mutations associated with macrolide resistance in clinical isolate CDC-SF003. Further, phylogenetics revealed that this strain belonged to the SS14 lineage, which correlated with its enhanced CDC typing method (ECDCT) strain type, 4d9f, as previously reported (7). While this 204 enrichment method yielded good results with isolates, most clinical specimens collected in this study had lower than 100 polA DNA copies/µl of 205 T. pallidum leading us to consider an alternative method. 206

207 SWGA has been shown to be successful with other bacterial pathogens in metagenomic samples; however, it has not been investigated with T. pallidum (25-27). We observed that samples enriched by SWGA using multiple primer sets exhibited a 10,000-fold increase in polA copy 208 number, with Pal 9 and 11 producing the highest relative percent T. pallidum DNA at 29% and 31%, respectively. While we chose to work with 209 Pal 11 as the optimal set, Pal 9 could also be a good alternative for enriching syphilis specimens. Further testing using Pal 11 showed that the limit 210 211 of detection was increased when compared to the T. pallidum enrichment obtained with NEB+MDA, with significant increases in both polA copy 212 number and percent T. pallidum across the 10-fold dilution series. Coverage across the T. pallidum genome exceeded 95% at 5X read depth for all 213 diluted samples, apart from the 1:10,000 diluted samples. Interestingly, we observed that increasing the input 100-fold resulted in a significant 214 decrease in the presence of RNP post-enrichment. Our data shows that >14 T. pallidum polA copies/ μ l can generate at least 95% coverage at 5X 215 read depth with the Nichols strain, which translated well to the clinical specimens tested. While there was a decrease in coverage in one of the 216 clinical specimens at 94.44% with 5X read depth when compared to the 98.62% coverage at 5X read depth observed in the 1:100 diluted Nichols 217 isolates, this could be primarily due to the improved capabilities of the NovaSeq 6000 when compared to the MiSeq v2 (500 cycle) platform used 218 to sequence this clinical specimen. Another possible reason for the variation in coverage could be due to the lower T. pallidum input copy number 219 in the clinical specimens.

The genomes derived directly from the 5 clinical specimens using SWGA were phylogenetically associated with the representative lineages (either Nichols-like or SS14-like) and also provided high levels of within lineage strain resolution, which is ideal for effective tracking of various strains circulating within a geographical area and outbreak investigations. In addition, the NGS methods described here can be used for macrolide resistance marker detection. As observed with NEB+MDA enrichment, *in silico* azithromycin mutation detection performed on the SWGA enriched specimens matched the results obtained with a real-time PCR, indicating that all clinical specimens contained the A2058G

- 225 mutation. SWGA-based enrichment also enabled sequencing of specimens within the range of detection limits for real-time PCR assays,
- suggesting that our NGS workflow can be adapted for *T. pallidum* detection in metagenomic samples.
- 227 In terms of expense, both methods are cost-effective for enriching *T. pallidum* genomic DNA, and while SWGA is cheaper than
- 228 NEB+MDA, sequencing reagents are the true limiting factor for WGS. With the recent advancements in large-scale sequencing platforms, overall
- sequencing costs can be further reduced. While NovaSeq 6000 has a much higher potential for multiplex sequencing, our data shows compatibility
- 230 of these enrichments for both NovaSeq 6000 and MiSeq platforms.
- 231 While we successfully enriched *T. pallidum* whole genomes in clinical specimens, the success of SWGA is limited by the constraint on
- primer size, which may reduce the selectivity for the target genome. Phi29 functions best between 30-35°C, and ramp-down incubations have been
- shown as an effective means of utilizing larger primers with increased melting temperatures (26-29). To help alleviate the constraints on primer
- size, we utilized a thermostable phi29 mutant which has a much higher optimal temperature at 45°C (30) compared to the 30-35°C functional
- range of the phi29 polymerase (26-27). This higher optimal temperature permits the use of longer oligonucleotides to be used in the SWGA
- reaction, potentially increasing the selectivity for the *T. pallidum* genome. The phi29 mutant has also shown to be more efficient, with a 3-hour
- exhaustion time when compared to the 8-16 hours required for the wild-type phi29 (30).

Our results show that SWGA is more sensitive, less cumbersome, and a faster method for enriching clinical specimens when compared to NEB+MDA, allowing for WGS of metagenomic samples with very low numbers of *T. pallidum*. In addition, the sequencing data generated is of sufficient quality to enable phylogenetic analyses and detection of mutations associated with azithromycin resistance. While the NEB+MDA was unsuitable for the clinical specimens in this study, our data suggests that it can be used for samples exceeding 129 genomic copies/µl.

242 Materials and Methods.

243 Specimen collection, *T. pallidum* strains used for WGS, and real-time qPCR. Specimens used in this study were collected from men

244 presenting with lesions of primary or secondary syphilis to the Emory Infectious Diseases Clinic, Emory University Hospital Midtown (EUHM) in

245 Atlanta, GA and St Louis County STD Clinic (STLC) in St. Louis, MO (Table 1). Patients were diagnosed with syphilis based on clinical

presentation and serology testing. Fourteen swab specimens were collected in Aptima Multitest storage medium (Hologic, Inc., Marlborough, MA)

247 at Emory Infectious Diseases Clinic and 1 specimen at St. Louis County STD Clinic (Table 1). All specimens were stored at -80°C until shipment

248 on dry ice to the CDC. The *T. pallidum* Nichols reference strain was used for initial optimization and validation of the two enrichment methods. A

recent rabbit propagated isolate, CDC-SF003, was also included for testing (Table 1; 7). Prior to study commencement, local IRB approvals were

obtained from, Emory University, and St. Louis County Department of Public Health, and the project was approved at CDC

251 DNA was extracted from specimens and rabbit testis extracts using the QIA amp DNA Mini Kit (Qiagen, Germantown, MD) following the 252 manufacturer's recommendations. Large-scale DNA extraction of three specimens was carried out on 1.5 ml of the Aptima stored specimen using 253 the QIA amp DNA Mini Kit following the manufacturer's recommendations for upscaling with slight modifications (Table 1). Proteinase K was 254 added at 0.1X total sample volume, and AL Buffer and absolute ethanol were added at 1X total sample volume. Each sample was processed through a single column, washed following the manufacturer's recommendations, and eluted in 100 µl AE Buffer (Qiagen). Following DNA 255 extraction, each sample was tested by a real-time quantitative duplex PCR (qPCR) targeting the *polA* gene of *T. pallidum* and human RNase P 256 gene (RNP) using a Rotor-Gene 6000 instrument (Qiagen) as previously described with modifications (7; see additional methods in supplemental 257 258 materials).

259	Enrichment of T. pallidum by capture of CpG methylated host DNA and multiple displacement amplification (MDA). Initially, DNA
260	concentration of extracts from clinical specimens and rabbit propagated strains were measured using the Qubit dsDNA HS assay (Thermo Fisher
261	Scientific, Waltham, MA). Capture and removal of CpG methylated host DNA from samples were carried out using the NEBNext Microbiome
262	DNA Enrichment Kit following the manufacturer's recommendations with modifications (New England Biolabs, Ipswich, MA). For all samples
263	tested, 250 ng of DNA was subjected to two rounds of bead capture using the NEBNext Microbiome DNA Enrichment Kit and enriched
264	treponemal genomic DNA was purified using AMPure XP beads (Beckman Coulter, Indianapolis, IN). Enriched DNA samples were stored at -
265	20°C until MDA was performed. MDA was carried out using the REPLI-g Single Cell Kit following the manufacturer's recommendations with
266	slight modifications (Qiagen). Each MDA reaction was incubated at 30°C for 16 hr. Following amplification, the polymerase was inactivated at
267	65°C for 10 min, samples were purified with AMPure XP beads, and eluted with 100 µl 1X AE Buffer (Qiagen). For each enrichment using the
268	REPLI-g Single Cell Kit, non-template controls were included to confirm the absence of <i>T. pallidum</i> .
269	A 10-fold dilution series on the Nichols strain was used to determine the limit of detection (LoD) for enrichment (see supplemental
270	materials) with NEB+MDA followed by sequencing on an Illumina NovaSeq 6000. After DNA extraction, each dilution in the series was enriched
271	by NEB+MDA, genomic copy numbers estimated by <i>polA</i> qPCR, and sequencing performed in triplicate. Enriched samples were diluted 1:10
272	prior to measuring RNP amplification. The LoD was set at the minimal genome copy number required to generate a \geq 5X read depth with \geq 95%
273	genome coverage compared to the reference genome.
274	Selective whole genome amplification (SWGA) primer design, validation, and enrichment. Primers with an increased affinity to T. pallidum
275	were identified using the swga Toolkit as previously described with slight modifications (https://www.github.com/eclarke/swga; 26; see
276	supplemental materials). Eight primer sets (SWGA Pal 1-8), including 4 additional primer sets (SWGA Pal 9-12) generated by combining primers

in the initial set (Table S1), were chosen for SWGA using the EquiPhi29 DNA Polymerase (Thermo Fisher Scientific, Waltham, MA). To account
for the 3'-5' exonuclease activity of the phi29 polymerase, all SWGA primers were generated with phosphorothioate bonds between the last two
nucleotides at the 3' end (Table S1). Each of the 12 primer sets were tested in triplicate against the spiked sample diluted to an estimated 100 *T*. *pallidum polA* copies/µl (see supplemental materials).

281 Prior to SWGA enrichment, samples were denatured for 5 min at 95°C by adding 2.5 µl of DNA to 2.5 µl denaturing solution, containing

custom primers, then placed immediately on ice until the Equiphi29 master mix, prepared as per manufacturer's recommendations, was added

283 (Thermo Fisher Scientific, Waltham, MA). MDA was carried out following the manufacturer's recommendations with modifications (Thermo

Fisher Scientific; 30). The reaction contained EquiPhi29 master mix, with EquiPhi29 Reaction Buffer at a final concentration of 1X, each primer

at a final concentration of 4 µM, and nuclease-free H₂O was added to a final reaction volume of 20 µl. Reaction tubes were gently mixed by pulse

vortexing and incubated at 45°C for 3 hr. MDA was stopped by inactivating the DNA polymerase at 65°C for 15 min. All reactions were purified

using AMPure XP beads and eluted in 100 μ l AE buffer (Qiagen). Non-template controls were included to confirm the absence of contaminate *T*.

288 *pallidum* DNA.

Relative percent *T. pallidum* in each sample was calculated as shown in Figure S1. SWGA Pal 11 was chosen for testing the LoD for downstream genome sequencing post-SWGA enrichment using the 10-fold dilution series, excluding the undiluted (neat) spiked sample. All enriched samples were validated by *polA* real-time qPCR in triplicate.

Sequencing and genome analysis of *T. pallidum* strains. Libraries were prepared using the NEBNext Ultra DNA Library Preparation Kit for
 NovaSeq and NEBNext Ultra II FS DNA Library Preparation Kit for MiSeq sequencing following the manufacturer's recommendations (New
 England Biolabs, Ipswich, MA). For the validation experiments, sequencing was carried out on the Nichols reference strain using the Illumina

NovaSeq 6000 platform following the manufacturer's recommendations (Illumina, San Diego, CA). Sequencing of isolate CDC-SF003 and swab
 specimens were carried out using the MiSeq v2 (500 cycle) platform following the manufacturer's recommendations (Illumina, San Diego, CA).

297 Post sequencing, reads were deduplicated, trimmed, and down selected for *T. pallidum* (supplemental materials). All down selected *T*.

298 pallidum reads were mapped to the T. pallidum reference genomes, and de novo assembled. Phylogenetic analyses were performed as described in

the supplemental materials. Apart from the genomes sequenced in this study, 122 high quality (with at least 5x read depth covering > 90% of the

300 genome) T. pallidum genomes deposited in the NCBI's Sequencing Read Archive (SRA) under the BioProject number PRJEB20795 and

301 PRJNA508872 were also included (12, 14). The publicly available raw sequencing data were re-analyzed to determine the quality as described in

302 the supplemental materials. A second phylogenetic tree was also reconstructed by including all the genomes sequenced from the 10-fold dilution

303 series for both NEB+MDA and SWGA enriched samples. Genomic sequencing data from samples included in the phylogenetic analyses covered

at least 90% of the reference genome with 5X read depth. Variant calls for the A2058G and A2059G macrolide resistance mutations were

305 validated using a real-time PCR assay as previously described (31).

306 Statistical analyses. Statistical analyses were performed in R (R Foundation for Statistical Computing, Vienna, Austria) using the R companion

307 software RStudio (Rstudio, Boston, MA). Statistical significance was determined by analysis of variance (ANOVA) and Tukey post hoc multiple

308 comparisons tests. T. pallidum percent DNA were normalized through Log_{10} conversions. Quantitative data are presented as means \pm standard

309	error.	Differences	were considered	statistically	/ signif	icant if a	P < 0.05.
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Data availability. All sequencing data associated with this study were submitted to the National Center for Biotechnology Information's sequence

read archive (SRA) under the BioProject accession ID PRJNA744275.

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318	Author Contributions
319	Allan Pillay and Ellen N. Kersh conceived the study. Allan Pillay, Charles M. Thurlow, Cheng Chen, and Lilia Ganova-Raeva designed
320	the study. Charles M. Thurlow and Allan Pillay designed the enrichment protocols. Charles M. Thurlow designed the SWGA specific custom
321	primer sets used during this study and performed all enrichment experiments. Charles M. Thurlow, Allan Pillay, Samantha S. Katz, Lara Pereira,
322	Alyssa Debra, Kendra Vilfort, Yongcheng Sun, Kai-Hua Chi, and Damien Danavall performed the laboratory experiments and assisted with
323	specimen collection. Kimberly Workowski, Stephanie E. Cohen, Hilary Reno, and Susan S. Philip collected clinical specimens and patient data.
324	Mark Burroughs, Mili Sheth, and Charles M. Thurlow performed Illumina sequencing. Sandeep J. Joseph performed the bioinformatic analyses of
325	the genomic data, phylogenetic analysis and contributed to the generation of tables and figures. Charles M. Thurlow and Sandeep J. Joseph
326	performed data analysis. Charles M. Thurlow wrote and prepared the manuscript with oversight by Allan Pillay and contributions from Sandeep J.
327	Joseph and Weiping Cao, which was reviewed by all authors for revisions.

328 Disclaimer

The findings and conclusions in this report are those of the authors and do not necessarily represent the official position of the Centers for
 Disease Control and Prevention. We declare that there are no competing interests.

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- 411
- 412 Tables

Sample/ Isolate ID	Collection Year	Source	Gender	Sexual Status	Syphilis Stage	Site of Lesion	Antibody Titer (Assay)	qPCR (<i>T. pallidum polA</i> in DNA Extract)	Extraction Method	Reference
CDC-SF003	2017	San Francisco	Male	MSM	Primary	Penis	1:4 (VDRL)	9, 680 copies/µl	Standard	Pereira et al., 2020
EUHM-001	2019	Atlanta	Male	MSM	Secondary	Neck	1:128 (RPR)	< 1 copy/µl	Standard	This study
EUHM-002	2019	Atlanta	Male	MSM	Secondary	Perianal	1:256 (RPR)	< 1 copy/µl	Standard	This study
EUHM-003	2019	Atlanta	Male	MSM	Secondary	Penis	1:32 (RPR)	< 1 copy/µl	Standard	This study
EUHM-004	2019	Atlanta	Male	MSM	Primary	Penis	1:4 (RPR)	106.7 ± 6.5 copies/µl	Standard	This study
EUHM-005	2019	Atlanta	Male	MSM	Secondary	Penis	1:64 (RPR)	< 1 copy/µl	Standard	This study
EUHM-006	2019	Atlanta	Male	MSM	Primary	Penis	1:16 (RPR)	< 1 copy/µl	Standard	This study
EUHM-007	2019	Atlanta	Male	MSM	Secondary	Hand	1:64 (RPR)	< 1 copy/µl	Standard	This study
EUHM-008	2019	Atlanta	Male	MSM	Secondary	Scrotum	1:64 (RPR)	$0.9\pm0.1~copy/\mu l$	Standard	This study
EUHM-009	2019	Atlanta	Male	MSM	Secondary	Scrotum	1:64 (RPR)	< 1 copy/µl	Standard	This study
EUHM-010	2019	Atlanta	Male	MSM	Secondary	Scrotum	1:128 (RPR)	< 1 copy/µl	Standard	This study
EUHM-011	2019	Atlanta	Male	MSM	Primary	Penis	1:32 (RPR)	< 1 copy/µl	Standard	This study
EUHM-012	2019	Atlanta	Male	MSM	Primary	Penis	1:8 (RPR)	$31.5\pm0.5\ copies/\mu l$	Large Scale	This study
EUHM-013	2020	Atlanta	Male	MSM	Secondary	Penis	1:64 (RPR)	$122 \pm 1.2 \; copies/\mu l$	Large Scale	This study
EUHM-014	2020	Atlanta	Male	MSM	Secondary	NA*	1:16 (RPR)	$103\pm6.7\ copies/\mu l$	Large Scale	This study
STLC-001	2020	St. Louis	Male	MSW	Primary	Penis	NR** (RPR)	$28.8\pm3.1\ copies/\mu l$	Standard	This study

Table 1. Clinical and laboratory data for specimens and clinical isolate CDC-SF003.

* Not available ** Non-reactive

Sample	Enrichment method*	Clonal complex	<i>T.pallidum polA</i> post enrichment genome copies/µl	Raw read pairs	Non-host read pairs	Total read pairs after QC	Read pairs classified as <i>T.</i> <i>pallidum</i>	Percent of total read pairs classified as <i>T.</i> <i>pallidum</i>	Mean read depth	Percent genome covered ≥1X	Percent genome covered ≥5X	Percent genome covered ≥10X
Nichols_CDC	non-enriched	Nichols-like	NA***	4,053,500	3,645,649	3,588,414	70,299	1.96	6.33	86.26	60.30	22.28
Nichols_CDC**	SWGA	Nichols-like	11,565,333 ± 1,294,672	3,701,303	3,692,932	3,648,044	3,414,111	93.59	751.17	98.39	98.24	98.16
CDC-SF003	NEB + MDA	SS14-like	2,394,930 ± 135,210	5,798,777	3,988,173	3,949,036	129,998	3.29	46.44	98.87	98.60	98.01
EUHM-004	SWGA	Nichols-like	6,367,089.5 ±240,811.5	6,102,826	4,440,618	4,280,401	1,403,645	32.79	370.39	96.99	95.13	92.67
EUHM-012	SWGA	Nichols-like	2,140,753 ± 28,192	10,350,274	5,870,287	5,716,082	2,793,693	48.87	639.86	96.34	93.98	91.89
EUHM-013	SWGA	SS14-like	5,159,716 ± 220,318.5	11,975,324	11,966,460	11,838,431	8,308,234	70.18	2,503.96	98.72	98.56	98.37
EUHM-014	SWGA	Nichols-like	2,573,508 ± 221,900.5	11,250,518	9,266,926	9,059,022	2,355,426	26.00	930.87	98.79	98.49	98.04
STLC-001	SWGA	SS14-like	$7,420,534 \pm 719,765$	11,293,960	7,770,834	7,721,767	3,004,631	38.91	1,133.43	98.32	95.94	94.10

Table 2. Sequencing percent coverage for the Nichols isolates, clinical isolate CDC-SF003, and clinical specimens across the T. pallidum reference genome.

^{*}All sequencing was performed using Illumina's MiSeq v2 (500 cycle) platform ^{**} Enrichment performed on 1,000 copies/µl *T. pallidum polA* input ^{***} Not available

414 Figures

415 Fig 1. T. pallidum polA copies/µl for the 10-fold dilution series spiked samples enriched by the NEBNext

416 Microbiome Enrichment Kit with REPLIg Single Cell MDA (NEB+MDA) or SWGA. The input *T*.

417 *pallidum polA* copies/ μ l for each dilution is displayed as Non-Enriched. The y-axis has been log_{10} scaled

- 418 fourdepired of it the Non-Enriched iditation series of Enriched iditation is a US Government work. It is not subject to copyright under 17 USC 105 and is also made available for use under 19 replicate enriched *T. pallidum* samples.
- 419 Teplicate enficienci *I. puttuum* samples.
- 420 Fig 2. Relative percent *T. pallidum* Nichols DNA for Non-Enriched, NEBNext Microbiome Enrichment
- 421 Kit with REPLIg Single Cell MDA (NEB+MDA), and SWGA enriched samples. Percent T. pallidum

422 DNA was calculated based on the input DNA concentration and *polA* copies/µl (Non-Enriched), and the

- 423 DNA concentration and *polA* copies/µl for the Nichols -spiked samples post-enrichment (NEB+MDA or
- 424 SWGA). The y-axis has been log₁₀ scaled for depiction of the Non-Enriched dilution series. Error bars
- 425 represent standard error among three replicate samples.
- 426 Fig 3. Percent coverage of sequencing reads of enriched *T. pallidum* Nichols spiked samples. (A)
- 427 Sequencing reads of samples enriched using the NEB Microbiome Enrichment Kit and REPLIg Single
- 428 Cell MDA (NEB+MDA). (B) Sequencing reads of samples enriched using SWGA. All samples were
- 429 sequenced using the Illumina NovaSeq 6000 platform. Error bars represent standard error between the
- 430 mapped reads derived from three replicate enriched Nichols samples.
- **Fig 4.** Percent coverage of isolates and clinical specimens. All samples were sequenced using the Illumina
- 432 MiSeq v2 (500 cycle) platform. Percent of *T. pallidum* reads are derived from down selected *T. pallidum*
- 433 reads. Prefiltered reads for Nichols-CDC were mapped to the Nichols reference genome (*NC_000919.1*).
- 434 The prefiltered reads in all clinical isolates and specimens were mapped against the SS14 reference
- 435 genome (*NC_021508.1*).
- **Fig 5.** SWGA primer set validation. (A) *T. pallidum polA* copies/µl for the Nichols mock sample (1:100
- 437 diluted) enriched with each SWGA primer set. (B) Relative percent *T. pallidum* DNA for the Nichols

- 438 spiked sample (1:100 dilution) enriched with each SWGA primer set. Percent T. pallidum DNA was
- 439 calculated based on the input DNA concentration and *polA* copies/µl for the Nichols mock samples post-
- 440 SWGA enrichment. The y-axis has been log₁₀ scaled for depiction of the relative percent *T. pallidum*
- 441 post-enrichment with each primer set. Error bars represent standard error among three replicate Nichols
- 442 samples. bioRxiv preprint doi: https://doi.org/10.1101/2021.07.09.451864; this version posted July 10, 2021. The copyright holder for this preprint (which was not c author/funder. This article is a US Government work. It is not subject to copyright under 17 USC 105 and is also made available for use under
- 443 Fig 6. Maximum likelihood global phylogenetic tree of the clinical isolate/specimen genome sequenced in
- this study along with publicly available *T. pallidum* genomes. The two major lineages, Nichols-like and
- 445 SS14-like are highlighted along with presence of genotypic mutation responsible for macrolide resistance
- and country of origin.

447



Input T. pallidum Dilution Factor















SWGA Primer Set



