Performance and automation of ancient DNA capture with RNA hyRAD probes

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Abstract
DNA hybridization-capture techniques allow researchers to focus their sequencing efforts on preselected genomic regions. This feature is especially useful when analysing ancient DNA (aDNA) extracts, which are often dominated by exogenous environmental sources. Here, we assessed, for the first time, the performance of hyRAD as an inexpensive and design-free alternative to commercial capture protocols to obtain authentic aDNA data from osseous remains. HyRAD relies on double enzymatic restriction of fresh DNA extracts to produce RNA probes that cover only a fraction of
In the last 15 years, high-throughput DNA sequencing (HTS) has found many applications in the genetic characterization of present-day biodiversity. HTS has also been extensively used to characterize past plant and animal communities, utilizing ancient DNA (aDNA) preserved both in sediments and in subfossilized tissues (Leonardi et al., 2016; Orlando & Cooper, 2014). For example, aDNA metabarcoding has helped identify ecological shifts in relation to climatic change (Pedersen et al., 2016; Willerslev et al., 2014) and human activities, including pastoralism (Giguët-Covex et al., 2014), despite limitations due to the extensively degraded nature of aDNA (Dabney et al., 2013). While massive parallel sequencing of short DNA fragments (Goodwin et al., 2016) has unveiled the full genome sequence of extinct organisms (even beyond the 1 million time range; van der Valk et al., 2021), aDNA libraries are frequently dominated by environmental microbial sources (Prüfer et al., 2010). Therefore, shotgun sequencing often provides a cost-ineffective strategy for characterizing past sequence variation at the genome scale.

In contrast to shotgun-sequencing, genome reduction approaches (sensu McCormack et al., 2013), including hybridization-based target-enrichment techniques (Kozarewa et al., 2015; Momanova et al., 2010), are designed to focus sequencing efforts on a fraction of the genome only. These approaches not only reduce analytical costs but also maximize the chances of identifying DNA present even in limited abundance (Slon et al., 2017). Therefore, target-enrichment provides an increasingly popular strategy for characterizing past biodiversity at both the community (Slon et al., 2017) and the population (Mathieson et al., 2015) levels. The approach is also increasingly used for genetic monitoring of endangered populations for which only noninvasive material can be collected from very limited and/or heavily contaminated sources (Aylward et al., 2018; Fontisere et al., 2020).

One common limitation of hybridization-based genome reduction techniques pertains to probe design and synthesis. Typically, nucleic acid probes are designed on the basis of existing molecular panels, which are, however, not available across all taxa, especially amongst nonmodel organisms. Probe synthesis also entails significant costs that are generally commensurate with the number of loci targeted. A number of in-house procedures have been developed to limit probe production costs, through PCR amplification of a number of target loci (Maricic et al., 2010; Peñaiba et al., 2014), or in vitro transcription when aiming at characterizing the whole genome (Carpenter et al., 2013). HyRAD technologies have been recently proposed to fill the gap between those two alternatives, allowing users to scale probe production to their research question and sequencing capacity (Schmid et al., 2017; Suchan et al., 2016). The methodology utilizes the versatility of RAD sequencing, which targets genomic regions flanking (Peterson et al., 2012) or encompassing (Baird et al., 2008) user-selected restriction sites. In hyRAD, enzymatic restriction is first applied to fresh DNA from a set of individuals of the focal or closely related species. Digested DNA fragments are then immortalized to produce target enrichment probes on-demand for in-solution aDNA capture (Suchan et al., 2016).

Despite reported success on plant and animal museum specimens (Boucher et al., 2016; Crates et al., 2019; Gauthier et al., 2020; Lang et al., 2020; Linck et al., 2017; Schmid et al., 2018; Toussaint et al., 2021), the full potential of hyRAD to characterize past molecular diversity remains largely unexplored. Currently, only one study retrieved aDNA over the 1,000-year time scale from ~7,000-year-old pine needles that were preserved in the lake sediments (Schmid et al., 2017). The suitability of hyRAD to (i) other preservation conditions,
(ii) deeper time ranges and (iii) calcified material, such as bones, teeth and shells, which represent the dominant fraction of the fossil record is, thus, unknown.

In this study, we applied hyRAD for the first time to ancient osseous material going beyond the radiocarbon time range and spanning diverse environmental conditions (from Tunisia, Poland and Russia; Table 1). We benchmarked hyRAD protocols including one or two rounds of capture and different combinations of restriction enzymes. These included methylation-sensitive enzymes with the aim of targeting hypomethylated genomic regions and limit the impact of post-mortem DNA damage (Seguin-Orlando et al., 2015; Smith et al., 2015), while diverting sequencing efforts from the most repetitive, hypermethylated, fraction of the genome (Karam et al., 2015; Larsson et al., 2013). We also developed and validated an automated protocol for targeted capture on Opentrons OT-2 liquid-handling robots to minimize hands-on time and the risk of experimental errors. Overall, the procedures presented in this study successfully retrieved authentic aDNA data from material beyond the radiocarbon time range and could achieve up to 146.46-fold enrichment of on-target reads on DNA extracts originally showing 0.34% endogenous DNA. The approach was successful in retrieving orthologous DNA fragments from different samples, even from limited sequencing efforts.

2 | METHODS

2.1 | DNA extraction and sample library preparation

The samples consisted of 12 bone or tooth samples, ranging from the Late Pleistocene to the 4th century CE (Common Era), from Tunisia, Poland and Russia. They were selected following shallow shotgun sequencing to encompass almost an entire range of endogenous DNA content (0.34%–73.78%; Table 1). DNA extraction followed Gamba et al. (2016) with modifications from Fages et al. (2019). Extracted DNA was treated with USER mix (Uracil-Specific Excision Reagent, New England Biolabs [NEB]) prior to DNA library construction, following Fages et al. (2019). A total of 14.9 μl of USER-treated aDNA extract was used for library preparation according to the modified Meyer and Kircher (2010) protocol presented in Fages et al. (2019), including one 6-bp external index (Orlando et al., 2013) and two 7-bp internal indices from Rohland et al. (2015) (Figure S1a). To ensure sufficient amounts of DNA templates for hyRAD capture, each of the 12 DNA libraries was amplified in two successive rounds of PCR. First, 3 μl of library was used within a 25-μl reaction with 0.2 μM of IS4 primer, 0.2 μM of indexing primer (Meyer & Kircher, 2010), 0.5 μg/μl of BSA (NEB), 1 μl of AccuPrime PfX polymerase (ThermoFisher Scientific) and 1× AccuPrime buffer. PCR consisted of 5 min denaturation at 95°C, followed by 11 cycles of 15 s at 95°C, 30 s at 60°C, 30 s at 68°C, and a final elongation for 5 min at 68°C. The number of PCR cycles for the second reaction was determined using qPCR and the samples were re-amplified either in two (samples KB2017, PLMie3, LOG3, DIV9, SV2019-22, SV2019-18, PLKaz4, PLKaz1; used for the capture with PstI–MseI probes only) or 16 parallel PCRs (samples PLMie10, PLSla2, PLMie8, PLKaz2; used for the capture with all three types of probes and the automated capture test, see below), to obtain the required amount for capture. The second amplification round was carried out using the ISS_reamp.P5 and ISS_reamp.P7 primers (Table S1) from Meyer and Kircher (2010) and the same PCR conditions as above, except that the primer concentration was 0.4 μM and only 1 μl of DNA template was used. For each sample, PCR replicates were pooled, purified and concentrated into 10 μl (for duplicate reactions) or 80 μl (for multiplicate reactions), using MinElute columns (Qiagen).

2.2 | Probe library preparation and probe production

Probe production was based on previous hyRAD protocols (Schmid et al., 2017), which rely on in vitro transcription of the probe library molecules containing T7 RNA polymerase promoter. Overall, high-molecular-weight DNA extracts (2 μg) from a modern domesticated horse were subject to double enzymatic restriction before ligation to two adapters showing terminal ends complementary to one restriction site (Figure S1b). Different combinations of restriction enzymes were selected to provide different levels of genome reduction. The PstI–MseI combination targeted 6-bp- and 4-bp-long restriction sites, respectively, and therefore was expected to produce fewer digested fragments than MspI–MseI and HpaII–MseI combinations, which both build on 4-bp-long restriction sites only. The latter two combinations were also selected to target genomic regions showing different DNA methylation background. Both HpaII and MspI enzymes target the same DNA restriction site (CICGG) but HpaII does not cleave DNA when the central CpG dinucleotide is methylated, in contrast to MspI.

Each digestion reaction used 40 μl of PstI–HF (or MspI/HpaII, NEB) and 20 μl of MseI (NEB) in the CutSmart buffer for 3 hr at 37°C in 100 μl. The reaction was purified using AMPure beads (Beckman Coulter), with a bead-to-liquid ratio of 2:1 and eluted in 20 μl of Tris 10 mm. The purified DNA was then used in a 3-hr ligation reaction at 16°C with 800 U of T4 DNA ligase (NEB) and 0.7 μM of each adapter (P1 and P2, see Table S1), with a final volume of 28 μl. Following ligation and purification with AMPure beads (bead-to-liquid ratio = 1.5:1), DNA templates were selected within the 190–390-bp size range using the Blue Pippin instrument and 2% agarose cassettes with external marker (Sage Science), which corresponds to inserts of ~100–300 bp. A further purification step allowed the elimination of those probe library constructs that did not incorporate the P2 adapter. This was achieved using the biotin group present in the P2 capture adapter and streptavidin-coated beads (MyOne C1 Dynabeads, Invitrogen). A total of 40 μl of the beads was first washed and resuspended in 2× TEN buffer (10 mM Tris-HCl, 1 mM EDTA, 2 mM NaCl), then combined with
| Sample name | Sample age | Site | Material | Endogenous DNA content (all/unique) |
|-------------|------------|------|----------|-----------------------------------|
| PLMie10     | Lusatian culture, Bronze Age/Early Iron Age (1,200–600 BCE) | Miechów, Poland | Metatarsus | 0.34/0.34% |
| PLSa2       | Funnelbeaker culture, Eneolithic (3,700–3,000 BCE) | Sławęcin, Poland | Molar | 0.51/0.51% |
| PLMie8      | Found in Funnelbeaker culture feature, Eneolithic (3,700–3,000 BCE) | Miechów, Poland | Metacarpus | 6.58/6.55% |
| DIV9        | Upper Palaeolithic, Late Glacial period (13,000–14,500 uncal BP) | Divnogor'ye, Voronezh Region, Russia | Molar | 27.42/1.72% |
| KB217       | Upper Palaeolithic | Medvezhiya cave, upper course of the river Pechora, Urals, Russia | Metapodium | 16.83/11.20% |
| PLMie3      | Przeworsk culture, Roman period (1st–4th century CE) | Miechów, Poland | Femur | 13.09/13.85% |
| LOG3        | Upper Palaeolithic (infinite 14C date, >52,200 uncal BP) | Hyena’s Lair, Altai Republic, Russia | Metatarsus | 20.64/20.55% |
| SV2019-22   | Iron Age, Middle Numidian period (5th–6th century BCE) | Althiburos, Tunisia | Tooth | 33.84/33.66% |
| SV2019-18   | Iron Age, Middle Numidian period (5th–6th century BCE) | Althiburos, Tunisia | Tooth | 47.19/46.97% |
| PLKaz4      | Trzciniec culture, Early Bronze Age (1,900–1,200 BCE) | Kazimierz Wielka, Poland | Petrous temporal bone | 51.67/51.48% |
| PLKaz1      | Trzciniec culture, Early Bronze Age (1,900–1,200 BCE) | Kazimierz Wielka, Poland | Petrous temporal bone | 60.43/60.15% |
| PLKaz2      | Trzciniec culture, Early Bronze Age (1,900–1,200 BCE) | Kazimierz Wielka, Poland | Petrous temporal bone | 73.78/73.59% |

Note: BCE = Before Common Era. CE = Common Era. BP = Before Present. Uncal = uncalibrated radiocarbon range.
40 µl of DNA, incubated rotating for 15 min, separated on a magnet, washed three times with 1× TEN buffer and resuspended in 15 µl of water. Finally, the probe library constructs showing both P1 and P2 adapters were PCR-amplified in a single reaction using KAPA HiFi HotStart ReadyMix (Roche), 0.6 µM of IS4 primer and indexing primer (Meyer & Kircher, 2010), and 7.5 µl of the bead solution obtained in the previous step. The PCR conditions were: 5 min denaturation at 95°C, followed by 14 cycles of 20 s at 98°C, 15 s at 60°C, 30 s at 72°C, and a final elongation for 5 min at 72°C. The probe library was purified using AMPure beads (bead-to-liquid ratio =1:1) and eluted in 20 µl of 10 mM Tris. Half of the probe library was kept for sequencing while the rest was digested with 1.5 U of MseI in a 15-µl reaction for 3 hr at 37°C to remove the P2 adapter prior to probe production through in vitro transcription. Adapter removal was confirmed and concentration estimated by running the sample on a TapeStation 4200 instrument (Agilent). RNA probes were then synthesized following in vitro transcription using the HiScribe T7 High Yield RNA Synthesis Kit (NEB) according to the manufacturer’s instructions, with 1/3 of UTP molarity as biotin-16-UTP (Roche), and 2 µl of the DNA template in a 40-µl reaction, for 16 hr at 37°C. Each reaction was then subject to TurboDNAse (Thermo Fisher) treatment at 37°C for 30 min to remove remaining DNA templates and purified on an RNEasy Mini column (Qiagen) using standard procedures, except that 675 µl of IS4 primer and indexing primer (Meyer & Kircher, 2010), and 7.5 µl of the bead solution after the capture and wash steps, and 0.5 µl of IS5_reamp.P5 and IS6_reamp.P7 primers (Table S1) from Meyer & Kircher (2010). All resulting sample and probe libraries were manually following two-round captures on four samples (PLMie10, PLSla2, PLMie8, PLKaz2) or two rounds of enrichment (all the samples). The four samples sequenced after the first and second round of capture were used to assess the impact of successive rounds of capture. Specifically, the sample libraries obtained following one round of capture were subject to either a second round of capture after re-amplification for ~500 ng of DNA, or to sequencing following fewer PCR amplification cycles. The number of PCR cycles was determined by qPCR and ranged from 12 to 20 for preparing the second round of capture, five to 15 for sequencing after the first round of capture and three or four for sequencing after the second. PCRs were performed using KAPA HiFi HotStart ReadyMix, 7.5 µl of the beads solution after the capture and wash steps, and 0.5 µM of IS5_reamp.P5 and IS6_reamp.P7 primers (Table S1) from Meyer and Kircher (2010). All resulting sample and probe libraries were sequenced on the Illumina MiniSeq instrument, using a 2 × 80 bp High-Output Kit.

The detailed protocol is available in the Supporting Information.

2.4 | Protocol automation

The automation protocol was implemented using Opentrons OT-2 robots (opentrons.com; also see May, 2019), equipped with magnetic and PCR modules, and includes all steps underlying hybridization capture, namely: (i) hybridization of aDNA libraries with the blocking mix, prewarming and addition of probes; (ii) incubation in the Opentrons PCR module; and (iii) prewarming of all washing buffers and performing washing by moving the samples to the Opentrons magnetic module for bead separation and buffer change, and back for incubation on the PCR module. The efficacy of the automated capture was assessed by comparison with the performance achieved manually following two-round captures on four samples (PLMie10, PLSla2, PLMie8 and PLKaz2).
2.5 | Data analysis

Paired-end DNA sequence data obtained from probe libraries were filtered and trimmed using cutadapt version 2.10 (Martin, 2011). We only kept those read pairs showing a fragment of the P1 adapter (consisting of the T7 promoter and the restriction cut-site) at the beginning of the first read, and the second cut-site at the beginning of the second read. For the P1 adapter, no more than three errors were allowed, including at best one base indel at read start. For the second adapter, full sequence match to the second cut-site and full overlap thereof were enforced. The resulting trimmed read pairs were then aligned against the horse EquCab3 reference genome (Kalbfleisch et al., 2018) using paleomix version 1.3.2 (Schubert et al., 2014) and both bwa version 0.7.17 backtrack algorithm (Li et al., 2009) and bowtie2 version 2.3.4.1 (Langmead & Salzberg, 2012). For bowtie2, we used the mapping parameters recommended by Poulet and Orlando (2020), while default parameters were used with bwa. All read alignments were then filtered to a minimal mapping quality of 25. While read alignment was carried out using both bwa and bowtie2 with similar outcomes, we chose the latter for subsequent analyses, based on generally higher numbers of mapped collapsed reads, in line with previous work (Cahill et al., 2018; Poulet & Orlando, 2020; Table S2; Figure S2).

For DNA sequences obtained from aDNA libraries, similar procedures were used, except that (i) adapterremoval version 2.3.1 (Schubert et al., 2016) was used for read trimming and collapsing instead of cutadapt; (ii) seeding was disabled during bwa mapping (Schubert et al. 2012); (iii) rmdup_collapsed from paleomix was used for removing PCR duplicates for collapsed reads and picard tools version 2.18.0 (http://broadinstitute.github.io/picard/) for uncollapsed reads; and (iv) post-mortem DNA damage was assessed using mapdamage version 2.2.1 (Jónsson et al., 2013) and pmdtools version 0.50 (Skoglund et al., 2014).

The expected number and size of DNA fragments obtained following enzymatic digestion were estimated in silico applying Fragmatic (Chafin et al., 2018) to the EquCab3.0 reference genome. Here, all three combinations of enzymatic restrictions could be investigated using two analyses only, as HpolI and MspI target the same restriction site. The preseq version 2.0.3 (Daley & Smith, 2013) functions c_curve (step size in extrapolations =10,000) and lc_extrap (maximum extrapolation =50,000,000, step size in extrapolations =10,000) were used to estimate library complexity profiles. This was applied to both probe sequence data and aDNA sequence data following one and two rounds of capture.

Comparison between the different experimental conditions was carried out by random sampling of identical numbers of reads for each given sample for all the treatments (to the lowest number of reads obtained for each sample (range = 88,013–690,062, mean = 464,500).

The following statistics were calculated for the libraries pre- and post-capture, using custom scripts based on samtools version 1.10 (Li et al., 2009) and bedtools version 2.29.2 (Quinlan & Hall, 2010): the fraction of endogenous DNA (number of reads mapping to the reference genome/number of raw reads); the fraction of unique endogenous DNA; the number of PCR duplicates; the fraction of on-target reads (reads showing at least 1-bp overlap with probe alignments); the fraction of unique on-target reads; the average depth-of-coverage for unique on-target reads; the size distribution of the uniquely mapped fragments (based on collapsed read pairs only, as these ensure that full aDNA fragments were sequenced); the GC content of the uniquely mapped reads; and the number of on-target sites with depth >0 (following PCR duplicate removal). Statistical significance was tested with Wilcoxon’s signed-rank-sum test in R (R Core Team, 2021), except for the differences between automatic and manual capture and statistics regarding damage rates of reads captured with methylation-sensitive probes, for which the sample size was too small. These latter analyses should therefore be considered as exploratory only. We have also plotted the size distribution of on-target reads (based only on collapsed reads for precision), and analysed enrichment folds (number of reads mapped after the procedure/number of reads mapped prior to the procedure) for captures with PstI–Msel probes. These data were further analysed using generalized linear models (GLMs) with inverse Gaussian distribution and the original endogenous content, level of PCR duplicates and age of the samples as independent variables. The relationship between enrichment-folds and read length was analysed using a GLM with Poisson distribution, stratified by sample, using the lme4 R package (Bates et al., 2015). For probe reads, additional statistics such as the %GC content, CpG content and fragment length were calculated. When probe paired-end reads were not overlapping, GC and CpG content were calculated based on the underlying genomic region on the reference genome.

To assess the authenticity of the obtained sequences, we quantified nucleotide misincorporations rates using mapdamage (Jönsson et al., 2013) and pmdtools (Skoglund et al., 2014). For data sets obtained using MspI–Msel and HpolI–Msel probes, we also used pmdtools to quantify CpG→TpG misincorporation rates. Such transitions generally derive from post-mortem cytosine deamination (Briggs et al., 2007). However, methylated CpG dinucleotides that are deaminated post-mortem are protected from USER-treatment and sequenced as TpG dinucleotides (Hanghøj et al., 2016), in contrast to unmethylated CpG dinucleotides that are converted into UpG dinucleotides and eliminated following USER-treatment (Hanghøj et al., 2016). Therefore, CpG→TpG misincorporation rates calculated on sequencing data generated following USER-treatment offer an opportunity to track the overall methylation levels of the genomic regions effectively enriched. We also assessed the fraction of mappable reads obtained using each probe set, and the proportion of reads flagged as multimapping to determine whether the targeted regions could represent genomic regions with nonaverage numbers of repeated elements.

For all 12 samples captured on PstI–Msel probes, genomic variants were called with freebayes version 1.2.0 (Garrison & Marth, 2012) using sequence data obtained precapture and after second round of capture, with the following options: --hwe-priors-off
HyRAD probes were found evenly distributed across all autosomes and the X chromosome (Figures S7 and S8), while no alignments were retrieved against Y-chromosomal contigs, in line with probes originating from a mare. Additionally, mitochondrial DNA sequences were extremely rare within the probe libraries (nine for HpaII, six for Mspl and absent for PstI; Table S3), as expected from the (nearly) absence of in silico-predicted digested fragments (absent for PstI–Msel and 10 for Mspl/HpaII–Msel).

### 3 | RESULTS

#### 3.1 | Study design and probe libraries

Over the last few years, our laboratory has undertaken an extensive genomic characterization of ancient horse specimens, which has resulted in an assessment of aDNA preservation levels in over 2,000 osseous remains, spread across the last 100,000 years in North Africa and Eurasia (Fages et al., 2019; Orlando, 2020). This allowed us to select 12 aDNA extracts representing three biomolecular preservation contexts and showing an entire range of endogenous DNA (from 0.34% to 73.78%; Table 1). Target-enrichment was applied using three panels of hyRAD probes prepared from the genomic DNA extract of a single modern horse that was digested by different combinations of restriction enzymes (PstI–Msel, Mspl–Msel and HpaII–Msel). The experimental procedures described here allowed for the preparation of extensive amounts of RNA probes. For instance, we obtained 92, 51 and 13.74 µg of RNA from 2 µl of digested PstI–Msel, Mspl–Msel and HpaII–Msel probe libraries, respectively. Digesting and using the full volume of probe libraries would, thus, theoretically allow users to perform ~400–2,700 capture reactions.

The size distribution of probe library templates was highly similar to that predicted after in silico digestion of the horse reference genome within the selected size range (Figure S4). From limited sequencing efforts (502,815 read pairs), we could estimate that 1.81% (PstI–Msel), 2.07% (Mspl–Msel) and 1.55% (HpaII–Msel) of the horse genome was represented in the probe libraries (Table 2). This equates to ~21%–29% (1.55%/7.16% and 2.07%/7.16%) to 38% (1.81%/4.77%) of the maximal genome coverage expected according to in silico digestion, as sequencing was limited. PRESEQ calculations (Figures S5 and S6) indicated that increasing sequencing efforts to ~5–10 million read pairs would be sufficient to reveal the full probe library content.

### Table 2 | Sequence characteristics for the sequenced probes libraries (*in-silico* calculations for HpaII–Msel probes do not take methylation-sensitivity into account)

| Enzyme combination | Raw reads | Properly paired reads | Number of multimapping properly paired reads | Number of targets after merging overlapping targets | Percentage of the genome targeted | Percentage of the genome predicted in silico |
|---------------------|-----------|-----------------------|----------------------------------------------|--------------------------------------------------|-----------------------------------|---------------------------------------------|
| PstI–Msel           | 502,815   | 380,495               | 43,713                                       | 247,716                                          | 1.81%                            | 4.77%                                       |
| Mspl–Msel           | 502,815   | 381,803               | 55,672                                       | 260,579                                          | 2.07%                            | 7.16%                                       |
| HpaII–Msel          | 502,815   | 387,275               | 32,569                                       | 234,924                                          | 1.55%                            | 7.16%*                                      |

HyRAD probes were found evenly distributed across all autosomes and the X chromosome (Figures S7 and S8), while no alignments were retrieved against Y-chromosomal contigs, in line with probes originating from a mare. Additionally, mitochondrial DNA sequences were extremely rare within the probe libraries (nine for HpaII, six for Mspl and absent for PstI; Table S3), as expected from the (nearly) absence of in silico-predicted digested fragments (absent for PstI–Msel and 10 for Mspl/HpaII–Msel).

#### 3.2 | Enrichment efficacy

Sequence alignments showed the hallmark of post-mortem DNA degradation following USER-treatment, including mild C→T misincorporation rates towards read termini (Figure S9) and fragmentation at cytosine residues (i.e., those that are deaminated into uracil after death and cleaved by USER-treatment; Rohland et al., 2015). This indicated that the experimental procedure followed in this study succeeded in retrieving authentic aDNA sequence data.

We first assessed hyRAD performance on four specimens by comparing the sequence data obtained following shotgun sequencing and after one or two successive rounds of capture (Figure 1). HyRAD capture resulted in increased overall levels of endogenous DNA present in the three aDNA libraries characterized by low starting endogenous DNA (0.34%–6.55% precapture). This was true for all three probe types, even though these only covered 4.77%–7.16% of the horse genome. For these three samples, the number of horse DNA fragments increased dramatically following one round of enrichment (2.73–21.63-fold for unique reads; 2.27–22.62-fold considering PCR duplicates). Endogenous DNA content increased even further following a second round of enrichment for the two samples (on PstI–Msel probes) or three samples (for the rest of the probes) with the lowest endogenous DNA content (4.00- to 53.13-fold and 4.17- to 73.07-fold, for unique reads and considering PCR duplicates, respectively). Endogenous DNA content was reduced after the second capture for the PLMie8 sample on PstI–Msel probes (6.55% original endogenous DNA content), from 3.10- to 2.64-fold for unique reads and 3.12- to 2.75-fold considering PCR duplicates. However, using the robotic protocol, the values after the second capture were higher (4.74- and 4.95-fold, respectively), pointing to the possible variability of the procedure outcome and benefits of
FIGURE 1 Percentage of endogenous, unique endogenous (after removing PCR duplicates), reads flagged as PCR duplicates, on-target reads (i.e., showing at least 1 nucleotide overlap with the target regions) and unique on-target reads. Results are shown for shotgun genomic libraries (pre) and the same libraries with one (capture 1) or two rounds (capture 2) of capture with three types of hyRAD probes, filtering for a minimum mapping quality of 25 (analyses relaxing the mapping quality filter are shown in Figure S10 so as to illustrate the impact of repeated elements in the sequence data).
FIGURE 2 Percentage of sites with non-null coverage, considering unique on-target reads. Results are shown for shotgun genomic libraries (pre) and the same libraries with one (capture 1) or two rounds (capture 2) of capture with the three types of hyRAD probes
duplication rate ($p = .0367$). Importantly, enrichment-fold levels increased with the fraction of large (>50-bp endogenous fragments available precapture (Figure 5; GLM, Poisson distribution, stratified by sample, $p < .0001$). This indicates that the prominent limiting factor for aDNA hyRAD capture is the extent of DNA fragmentation ongoing after death, and hence the availability of DNA templates of sufficient size during enrichment.

The increased on-target coverage achieved using hyRAD translated into a higher proportion of shared genomic loci amongst samples (Figure 6a). With our limited sequencing efforts, ~100,000 sites were common to four samples following shotgun sequencing and no sites were shared across all samples. With equivalent sequencing efforts, ~100,000 sites were common to 11 samples and 2,846 were shared across all samples. This led to the identification of 87 variants, 44 of which were present in at least two samples. In contrast, prior to capture, we only detected three variants that were private to two individual samples (Figure 6b,c). The total number of variants identified per sample also increased up to 52 following two rounds of hyRAD capture (Figure 6c). This demonstrates the capacity of hyRAD capture to provide shared orthologous sequence information across multiple samples. PRESEQ calculations showed that after the second round of capture, the number of unique horse reads present in DNA libraries could be discovered faster than after one unique round of capture (Figure 5c). As such reads are also significantly more frequently found on target after the second than the first round of capture (Figure 1), this indicates the second round of capture as enriching more efficiently for those loci of interest. It follows that researchers can achieve the sequencing of the shared variation within multiple orthologous regions across many samples with lower sequencing efforts.

**FIGURE 3** The effect of one (capture 1) or two rounds (capture 2) of capture with three types of hyRAD probes on the %GC content and sequenced fragment lengths, as compared with the shotgun libraries (pre)
Our preseq calculations can also provide guidance for making appropriate decisions on optimal sequencing efforts and, ultimately, scaling experimental costs. In particular, preseq calculations can predict how many reads mapping to the focal genome would be necessary before reaching saturation. Here, this number corresponds to a total of 20–50 million (paired-end) reads for the vast majority of samples (Figure S12). The only exception was sample DIV9, for which the full library content was discovered when ~10 million reads were sequenced, as the amount of endogenous DNA preserved was extremely limited. We therefore suggest that a conservative starting number of 20–50 million reads are first generated before running preseq calculations and committing to further sequencing costs.

### 3.3 Scaling and DNA methylation sensitivity

The choice of restriction enzymes allows hyRAD users to scale their experimental design according to the question of interest and sequencing capacity. By choosing restriction enzymes that are sensitive to DNA methylation (e.g., HpaII), they can also target less repetitive hypomethylated regions (Karam et al., 2015; Larsson et al., 2013). This may be especially interesting for aDNA characterization as such regions are less prone to post-mortem DNA damage (Smith et al. 2014, Seguin-Orlando et al., 2015). To demonstrate this, we compared on-target CpG→TpG misincorporation rates as a measure of post-mortem cytosine deamination, following one and two rounds of hyRAD capture with HpaII–MseI probes (methylation-sensitive).
and MspI–MseI probes (methylation-insensitive). These rates were indeed lower when the methylation-sensitive combination was used (Figure 7), but not when including off-target reads (Figure S13), confirming that the genomic fraction effectively enriched was associated with lower post-mortem DNA damage. The number of on-target reads flagged as multimapping was also considerably smaller when the methylation-sensitive combination was used (Figure S14; Table 2). This was not true for off-target reads, confirming that capturing hypomethylated regions can indeed help focus on genomic regions with reduced repetitive content.

3.4 | Protocol automation

Automated solutions to aDNA analyses have gained increasing interest over the last few years (Rohland et al., 2015; Slon et al., 2017), with the benefit of increased efficiency and reproducibility (Holland & Davies, 2020). However, investment costs for acquiring liquid-handling robots can be extremely prohibitive and not available outside of large-scale facilities. To provide an automation solution for most laboratories, we developed an automated hyRAD protocol that can be run on the inexpensive liquid-handling devices produced by Opentrons. Our automated implementation reduced hands-on time to 30 min per capture session, vs. 3 hr when carried out manually by skilled laboratory staff, while achieving similar performance (Figure 8). The developed protocol is available at https://github.com/TomaszSuchan/opentrons-hyRAD.

4 | DISCUSSION

In this study, we investigated for the first time the potential of hyRAD target-enrichment for aDNA extracted from osseous remains. First,
we modified the original hyRAD protocol in order to produce RNA probes from the ddRAD-seq templates, by including T7 polymerase promoter in the library adapter’s sequence, as in Schmid et al. (2017). This modification has several benefits over the original approach (Suchan et al., 2016). First, using RNA probes reduces the risk of contamination of aDNA libraries by the probes during capture. Second, RNA-DNA heteroduplexes show higher affinity than DNA-DNA homoduplexes (Lesnik & Freier, 1995), thus improving capture efficacy (Furtwängler et al., 2020). Finally, the T7 polymerase reaction allowed us to obtain sufficient amounts of RNA probes to carry out thousands of capture reactions from relatively modest amounts of starting DNA. We thus solved the main limitation of the original hyRAD protocol, which required many subsequent PCR amplifications to obtain sufficient amounts of capture probes (Suchan et al., 2016).

We assessed hyRAD performance using three types of probes and one single or two rounds of enrichment. We found that the second round of enrichment consistently improved on-target recovery of unique reads, despite also dramatically increasing the proportion of PCR duplicates. This was especially the case for samples of low endogenous DNA content, consistent with previous research (Fontsere et al., 2020; Hernandez-Rodriguez et al., 2018). We also confirmed previous reports of limited enrichment for aDNA libraries showing high starting endogenous DNA content (Cruz-Dávalos et al., 2017).

We show that this was mainly driven by the dramatic drop of mean mapping qualities for the captured reads, probably resulting from increased proportions of repeated elements post-capture.

On-target enrichment reached up to 146-fold for the sample with the lowest endogenous DNA content (0.34%) and around 3.5-fold for samples with 13.90% and 20.64% of endogenous DNA content (not counting samples with high initial PCR duplication rates), which reduces sequencing costs proportionally. Enrichment-fold values were, however, limited to around 1.4–1.8 for samples with >30% endogenous DNA (Figure 4; Table S5), which is lower than reported for commercial capture protocols (Cruz-Dávalos et al., 2017). Commercial protocols, however, are based on synthetic probes, and can be carefully designed to represent optimal probe molecular features that cannot be controlled with hyRAD. Our experimental conditions were also less stringent (55°C incubation for 40 and 16 hr during the first and second rounds of enrichment, respectively), in line with the recommendations of Cruz-Dávalos et al. (2017) on synthetic RNA probes. Further work is necessary to assess whether more stringent annealing conditions could overcome some of the limitations of our current procedure.

Our results confirm that libraries showing high initial rates of PCR duplicates following shotgun-sequencing (samples DIV9 and KB217 in our case) perform the poorest after enrichment. This demonstrates that library complexity levels have a strong impact on capture outcomes, but that our procedure still succeeds in retrieving substantial amounts of on-target reads even when library complexity is limited (Figure 4; Table S5).

The size distribution of horse DNA fragments increased following each round of enrichment, as previously shown for aDNA and environmental DNA (Cruz-Dávalos et al., 2017; Enk et al., 2014). This suggests that more limited enrichment success can be expected for aDNA libraries characterized by extreme fragmentation. Therefore, we suggest that the enrichment potential of a given set of samples can be assessed prior to capture following shallow shotgun sequencing so as to gauge for both complexity levels (i.e., low duplication rates) and the presence of endogenous DNA fragments of a relatively long size (e.g., 80 bp and above).
We also demonstrated that probe preparation with methylation-sensitive restriction enzymes could help focus sequencing efforts on hypomethylated regions, and limit both the fraction of the repetitive regions sequenced (Karam et al., 2015; Larsson et al., 2013) and the amounts of nucleotide misincorporations pertaining to post-mortem DNA damage (Smith et al. 2014, Seguin-Orlando et al., 2015). Therefore, the hyRAD procedure described can not only achieve a cost-effective characterization of orthologous genomic regions across a set of samples but can also improve the underlying sequence quality. Additionally, automation of the procedure can reduce hands-on time, and improve experimental reproducibility and traceability. Our approach appears especially appropriate for ancient samples characterized with low endogenous DNA content but not excessively fragmented. It may provide a future avenue for the genetic characterization of environmental (Wilcox et al., 2018) and noninvasive samples (e.g., faeces, Fontsere et al., 2020), which are both characterized by limited DNA amounts of the species of interest.

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AUTHOR CONTRIBUTIONS

Designed and conceived the study: T.S. and L.O.. Performed wet-laboratory work: T.S., M.K., N.K., L.C., L.T.C. and S.S. Provided samples, material and reagents: J.S., M.K., K.K., J.K., A.N.B., A.A.B., A.S.G., S.V.L., J.W., S.P., K.T., M.N., M.M.dH., A.A.T., A.J.E.P., A.K.O. Performed wet-laboratory work: T.S., M.K., N.K., L.C., L.T.C. and S.S. Provided samples, material and reagents: J.S., M.K., K.K., J.K., A.N.B., A.A.B., A.S.G., S.V.L., J.W., S.P., K.T., M.N., M.M.dH., A.A.T., A.J.E.P., A.K.O. and L.O. Analysed data: T.S. and L.O. Interpreted the data: T.S., L.O. and innovation programme (grant agreement 681605).

DATA AVAILABILITY STATEMENT

The data sets generated for this study can be accessed in the ENA (Accession No. PRJEB43744). The laboratory protocol is available at https://doi.org/10.17504/protocols.io.bywxpxfn. The program for piloting the Opentrons OT-2 robot can be accessed at https://github.com/TomaszSuchan/opentrons-hyRAD/tree/v1.0.

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