

Dominant components of the Thoroughbred metabolome characterised by 1H-NMR spectroscopy: a metabolite atlas of common biofluids

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- Dominant components of the Thoroughbred metabolome characterised by ¹H-NMR 1 2 spectroscopy: A metabolite atlas of common biofluids. 3 E. Escalona*¹, J. Leng², A. Dona¹, C. Merrifield², E. Holmes¹, C. Proudman³ and J. Swann² 4 ¹Section Computational and Systems Medicine, Division of Surgery and Cancer, Faculty of 5 6 Medicine, Imperial College London, Sir Alexander Fleming Building, London SW7 2AZ, 7 UK; 8 ²Department of Food and Nutritional Sciences, University of Reading, Whiteknights, 9 Reading, Berkshire, RG6 6AH, UK; ³ Department of Gastroenterology/School of Veterinary Medicine, University of Liverpool, 10 11 Neston CH64 7TE, UK. 12 13 *Corresponding author email: e.escalona10@imperial.ac.uk 14 **Keywords:** horse; metabonomics; metabolomics; metabolites; biofluids; nuclear magnetic 15 16 resonance 17 18 19 Summary
- 21 screening and investigating mammalian metabolism. This study aims to create a metabolic

Reasons for performing study: Metabonomics is emerging as a powerful tool for disease

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- 22 framework by producing a preliminary reference guide for the normal equine metabolic
- 23 milieu.
- 24 **Objectives:** To metabolically profile plasma, urine and faecal water from healthy racehorses
- 25 using high resolution ¹H-NMR spectroscopy and to provide a list of dominant metabolites
- 26 present in each biofluid for the benefit of future research in this area.
 - 27 **Study design:** This study was performed using seven Thoroughbreds in race training at a
 - 28 single time-point. Urine and faecal samples were collected non-invasively and plasma was
 - 29 obtained from samples taken for routine clinical chemistry purposes.
 - 30 **Methods:** Biofluids were analysed using ¹H-NMR spectroscopy. Metabolite assignment was
 - achieved *via* a range of 1D and 2D experiments.
 - 32 **Results:** A total of 102 metabolites were assigned across the three biological matrices. A core
- 33 metabonome of 14 metabolites was ubiquitous across all biofluids. All biological matrices
- provided a unique window on different aspects of systematic metabolism. Urine was the most
- 35 populated metabolite matrix with 65 identified metabolites, 39 of which were unique to this
- 36 biological compartment. A number of these were related to gut microbial host co-
- 37 metabolism. Faecal samples were the most metabolically variable between animals; acetate
- 38 was responsible for the majority (28%) of this variation. Short chain fatty acids were the
- 39 predominant features identified within this biofluid by ¹H-NMR spectroscopy.
- 40 **Conclusions:** Metabonomics provides a platform for investigating complex and dynamic
- 41 interactions between the host and its consortium of gut microbes and has the potential to
- 42 uncover markers for health and disease in a variety of biofluids. Inherent variation in faecal
- 43 extracts along with the relative abundance of microbial-mammalian metabolites in urine and
- 44 invasive nature of plasma sampling, infers that urine is the most appropriate biofluid for the
- 45 purposes of metabonomic analysis.

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Introduction

Metabonomics is a powerful systems biology approach that aims to simultaneously measure all the low molecular weight metabolites present in a biofluid or tissue. This approach to global untargeted characterisation of the metabolic phenotype allows the study of multidimensional biochemical responses of complex biological systems to genetic or environmental stimuli [1]. Metabolic profiling captures information from both intrinsic (genetics, protein expression) and environmental inputs (diet, gut microbiota), providing holistic information on the global system. This strategy has proven highly effective for unrayelling the complex metabolic interactions between the mammalian host and its resident gut microbiota. Metabonomics is a tool of particular interest to equine researchers given the vast impact of the equine gut microbiome on the bioavailability of food, medication and energy. Metabonomics, along with other 'omic' technologies such as genomics, proteomics and transcriptomics is increasingly showing potential in clinical settings as both a screening tool and a means for mechanistic elucidation of disease pathways [2-4]. To date, there are fewer metabonomic studies exploring veterinary concerns than there are human and rodent studies. The majority of mammalian work has concentrated on laboratory animals [5-7] and animal models with high translatability to human health such as the pig [8; 9]. Less attention has been given to herbivorous hind-gut fermenters and the majority of equine metabolic work concentrates on drug detection within the racing industry [10, 11]. Equine-specific metabonomic studies include the use of the horse as a model for the metabolic response to a dextrose challenge in type-2 diabetes [12] and a laminitic plasma NMR study using an oligofructose overload model [13]. Other equine studies include metabolic analysis of biofluids in response to age [14] and osteochondrosis [15]. Pappalardo

71 <i>et al</i> .	have explored metabolic variation in association with breed of horse and importantly
72 revea	aled significant differences that are likely to be attributed to differing growth rates and
73 prote	in utilisation [16]. These bodies of equine metabonomic work address the enormous
74 poten	ntial for exploring normal equine physiology and pathology-based variation. However,
75 there	is still a paucity of baseline data on the metabolic phenotype of horses and this study
76 serve	s as a reference tool for the Thoroughbred racehorse for clinicians wishing to use
77 metal	bonomic technologies to complement their research in either a diagnostic or mechanistic
78 capac	city.
79 Two	major analytical platforms are commonly used for metabolic profiling, nuclear magnetic
80 reson	nance (NMR) spectroscopy and mass spectrometry (MS). These approaches
81 simul	Itaneously capture quantitative information from a range of low molecular weight
82 metal	bolites across various different sample types. In metabonomic studies, NMR and MS are
83 used	both independently and in tandem to achieve an in-depth coverage of the metabolome.
84 In th	is study we comprehensively characterise the dominant features within the urinary,
85 plasm	na and faecal metabolomes of Thoroughbred racehorses using ¹ H-NMR spectroscopy, as
86 this p	provides a reliable, reproducible [17] screening tool and is non-destructive of samples.
87 Thord	oughbred racehorses represent a highly uniform equine population both genetically and
88 envir	onmentally. Normal clinical chemistry parameters exist for this equine subtype to assist
89 racing	g industry standards and this study will build upon data already in the literature and in

Materials and methods

Sample collection

clinical use.

From a large–scale study of urinary metabonomics of in-training Thoroughbred across a number of yards, 7 animals were selected at random to provide a baseline metabolic reference point for future comparison. Due to ethical constraints of invasive sampling and logistical limitations of simultaneous collection of all 3 biofluids, 7 animals were selected as representative of a wider population of animals. Early morning free-catch urine and faecal samples were collected into sterile plastic containers and snap-frozen in liquid nitrogen within 2 h. Plasma was obtained from excess clinical samples. Plasma samples were collected into heparinised tubes, spun down to obtain the plasma fraction and snap-frozen within 2 h of collection. All samples were then stored at -80°C prior to NMR analysis. Comprehensive metadata for each horse is shown in Supplementary Item 1 and samples were consistently taken before a morning concentrate feed.

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- 107 Sample preparation
- 108 Plasma and urine sample preparation was carried out as described by Beckonert et al.
- 109 2007[18].

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- 111 Faecal samples (100 mg) were combined with 1.7 mm Zirconia beads and 1 mL of distilled
- water and homogenised in a bead-beater for 10 min and centrifuged at 13,000 g for 10 min.
- Water was evaporated from the samples using vacuum concentrator (Speed-Vac) and then
- 114 reconstituted in 700 µL phosphate buffer (pH 7.4; 100% D₂O) containing 1 mM sodium 3-
- trimethylsilyl-1-[2,2,3,3-2H4]propionate (TSP).

- 117 Combined sampling approach for 2D NMR Experiments
- 2D spectra were obtained from pooled samples for each sample type to ensure comprehensive
- capture of metabolites. 50 µL of urine was pooled from each horse before the addition of 500

- $120~\mu L$ of phosphate buffer (and $500~\mu L$ of the resulting volume was added to the 5 mm NMR
- tube). The same process was repeated for the plasma samples with the addition of 500 μ L of
- 122 D₂O saline solution. For the faecal samples, individual samples were processed as previously
- described and then pooled after reconstitution with 100 μL of phosphate buffer.
- 124 Acquisition of 1D ¹H-NMR spectra
- 125 Spectroscopic analysis of all samples was carried out on a 700 MHz Bruker NMR
- 126 spectrometer^b operating at 300K and equipped with a 5 mm ¹H(¹³C/¹⁵N) inverse cryoprobe.
- 127 Urine and faecal samples
- 128 Standard one-dimensional ¹H-NMR spectra were acquired for all urine and faecal samples.
- We employed a standard one-dimensional pulse sequence (noesypr1d) that employs the first
- increment of a NOE sequence to achieve suppression of the water resonance with water peak
- 131 suppression using a standard pulse sequence [19]. For each sample, 8 dummy transients were
- followed by 256 transients and collected in 64K data point. Irradiation of the solvent (D₂O)
- resonance was applied during presaturation delay (2.0 s) for all spectra. The pulse sequence
- parameters including the 90° pulse, receiver gain and pulse powers were optimised for each
- sample set run. The spectral width was 20 ppm for all spectra. The free induction decay (FID)
- was processed with an exponential line broadening of 0.5 Hz prior to Fourier transformation.
- 137 Plasma samples
- Water-suppressed Carr-Purcell-Meiboom-Gill (CPMG) spin-echo spectra were acquired for
- the plasma samples, Here, 8 dummy scans followed by 256 scans were acquired for each
- sample in 64k data points with a total spin-spin relaxation delay of 1.5 s and a total delay
- between pulse cycles of 4.85 s.
- Acquisition of 2D ¹H-NMR spectroscopy was undertaken with an 800 MHz Bruker NMR
- spectrometer^b operating at 300K and equipped with a triple-resonance probe (TXI). J-

resolved spectroscopy (J-res) spectra were acquired from all biofluid composite samples using 64 transients per increment with 160 increments in the second dimension. The F1 (Jcoupling) domain spectral width covered 120 Hz. Prior to the double FT and magnitude calculation, the F1 data was zero-filled to 1024 points. The spectra were then tilted by 45° to provide orthogonality of the chemical shift and coupling constant axes and subsequently symmetrised about the F1 axis. ¹H-¹H Correlation Spectroscopy (COSY) was performed on all 3 types of pooled biofluid samples in order to detect correlations between protons on adjacent carbons. Transients were acquired with 4096 data points (sweep width of 7200 Hz in both axis) with 64 scans per increment and 320 increments in the F1 axis. The relaxation between successive pulse cycles was 2.3 s and were weighted using a sine bell function in T₁ and T₂ prior to fourier transformation and subsequently symmetrised about the diagonal axis. Two-dimensional echo/anti-echo ¹H-¹³C heteronuclear single quantum correlation (HSQC) spectra were also obtained. 256 scans were collected (16 dummy scans) at a spectral resolution of 4k in F2 across a spectral width of 12 ppm for ¹H and 170 ppm for the ¹³C axes. An acquisition time of 0.852 s and a relaxation delay of 1.2 s were used and delays were set for a 145 Hz one bond ¹H-¹³C coupling constant. Spectra were zero-filled in the F2 dimension by a factor of 2 to 8k, and zero-filling and linear prediction was applied in F1 to result in a resolution of 1k.

NMR spectral data pre-processing

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Data [-1.0 to 10.0 ppm] were imported into MatLab environment (7.0 The Mathsworks^c), where they were automatically phased, baseline-corrected and referenced to TSP (δ 0.00) or glucose (δ 5.233) for plasma using scripts written in-house [20]. To reduce analytical variation between samples the residual water signal (4.67 – 4.98 ppm) was truncated from the

data set. Probabilistic-quotient normalisation was used on each biofluid class separately to account for differing sample dilutions [21].

Data analysis of biological matrices

Unsupervised multivariate analysis was undertaken to visualise clustering and differences between samples [22]. Principal component analysis (PCA) was constructed using unit-variance scaled data (UV) [23]..

Metabolite assignment of endogenous metabolites was made by compiling the following information from each peak: the chemical shift and relative integral height from 1D spectra, the multiplicity using J-res spectra, the proton coupling information from COSY spectra and carbon shifts from HSQC spectra. This information was then used to search for matching metabolites from in-house databases, online databases (http://www.hmdb.ca/) and reference to published literature data [24-26]. Statistical Total Correlation Spectroscopy (STOCSY) was also employed to aid metabolite identification [27]. This method uses statistical connectivity between data points within a spectral profile. However, unlike 2D-NMR this method will also pick up metabolites involved in the same pathways due to biological covariance [28].

Results

Assignment of dominant metabolites across easily obtainable biofluids yields a preliminary equine metabolic atlas

The majority of known NMR-detectable metabolites were assigned in each biological matrix (Fig 1), with numerical reference to Table 1. Metabolite assignments were performed using two-dimensional correlation (COSY) and J-resolved (J-res) spectroscopy to ascertain peak

190 multiplicity, coupling constants and to overcome peak overlap. This is demonstrated in the 191 COSY (Fig 2A) and the J-res (Fig 2B). Heteronuclear Single Quantum Coherence (HSQC) was also performed to provide ¹³C shifts and confirm assignments. Putative metabolite 192 193 assignment was made using ¹H and ¹³C chemical shifts, peak multiplicity, coupling constants and relative peak integrals. Overall, 102 metabolites were identified by ¹H-NMR in the 3 194 195 biofluids (Table 1). Detailed assignment information is shown in Supplementary Item 2. 196 197 Cross-compartmental analysis revealed a core metabonome, along with compartment specific metabolites 198 199 Metabolites assigned to the 3 biofluids were compared to ascertain ubiquitous metabolites 200 and those that were specific to each biological compartment. The metabolic variation across 201 the biological matrices is displayed in a Venn diagram (Fig 3) to easily visualise inter-202 compartmental overlap. A total of 14 metabolites were ubiquitous to all biofluids, which we 203 will refer to as 'core' metabolites. These include energy-related metabolites such as glucose and lactate as well as a number of amino-acids including alanine, arginine, glycine, 204 205 glutamine, taurine, threonine and valine. The microbial related metabolites acetate, formate and p-hydroxyphenylacetate were also conserved across all biofluids studied. 206 207 PCA revealed that inter-animal metabolic variation was lower than the variation between the 208 different biofluids as visualised via clustering in the PCA scores plot (Fig 4A). As would be 209 expected, the PCA samples cluster based on biofluid type. However, importantly the degree 210 of clustering is different amongst biological matrices. Faecal samples demonstrate that they 211 are inherently variable compared to either matched urine or plasma samples. The faecal 212 metabonome displayed relatively higher concentrations of SCFA (butyrate, acetate and 213 propionate) compared to plasma and urine. Urine samples had relatively higher level of gutmicrobial co-metabolites such as hippurate, phenylacetylglycine (PAG), *p*-cresyl sulfate and trimethylamine-*N*-oxide (TMAO). In contrast, energy-related metabolites such as glucose and pyruvate were observed in relatively higher concentrations in plasma compared to urine. Variance plots shown in Fig 4A and B display the mean spectra of faeces and urine respectively coloured by the variance, represented as a percentage of the total variance. Here, creatinine can be seen to represent over 10% of the total variance seen within the urinary profiles (Fig 4B) and acetate accounted for over 28% of the total variance observed in the faecal profile (Fig 4A) compared to 1.8% in the plasma profile (not shown).

223 Plasma

Equine plasma contained relatively few metabolites compared to urine and faeces when analysed by NMR spectroscopy, partly as a result of the overlap of signals from macromolecular components such as lipoproteins and low molecular weight chemicals. A total of 38 metabolites were identified (Table 1), and of these 12 were unique to this biofluid, including the amino-acids tyrosine and phenylalanine and ketone bodies (α - and β -hydroxybutyrate). Plasma was observed to contain the highest levels of glucose compared to other biofluids within the data matrix.

231 Urine

The equine urinary metabolic profile was the most metabolically abundant of all biofluids measured (Table 1). In total 65 metabolites were identified. Of these, 11 were unique to urine and plasma, 3 were present in both urine and faeces and 39 metabolites were specific to urine. Urinary-specific metabolites included a number of aromatic compounds that arise from microbial-host co-metabolism. Hippurate, PAG, *p*-cresyl glucuronide and sulphate were

- 237 notably prominent in the aromatic region of the horse urine spectra (Fig 1B). Other urine specific metabolites of note include dietary compounds such as proline betaine.
- 239 Faeces
- 240 The faecal metabotype contained 43 metabolites, including 21 specific to faeces (Fig 3).
- 241 These metabolites include microbial fermentation products, SCFA (butyrate and propionate),
- and a number of dietary metabolites (caprylate and arabinose, maltose, glycerol and xylose).
- 243 Acetate was present in significantly greater concentrations than other metabolites.

Discussion

This is the first systematic description of the dominant metabolites of the healthy Thoroughbred racehorse and is important in establishing a metabolic reference from which to compare pathology-related variation. Characterisation of the equine metabolome highlights the diversity of information available in different biological matrices and as such provides a useful guide for researchers. Importantly, as has been shown in other species, urine provides a biological window into host-microbial metabolic interactions in the horse [8]. Although faeces may be considered a more direct representation of microbial metabolism, Fig 4A and B highlights the relatively greater variation in the faecal metabolic profiles compared to plasma and urine. Additionally, microbial-derived compounds absorbed from the gut are commonly not well represented in the faecal signature rendering the biological usefulness of this biofluid in gastrointestinal disease and mammalian-microbial co-metabolsim questionable.

A ubiquitous metabonome is present amongst mammals. The core equine metabonome comprises metabolites present in all 3 biofluids studied. 14 metabolites were identified including amino-acids, gut microbial metabolites and energy metabolites. Many of these

metabolites are ubiquitous, with varied functions and located in numerous tissue types. 261 262 Similar findings were reported by Merrifield et al. with 22 metabolites shared across 4 263 biological matrices in pigs [8]. 264 Marked ¹H-NMR metabolic similarities between horses and humans are suggested by observations of a number of shared metabolites. In total, there were 32 plasma, 23 urine and 265 266 27 faecal NMR detectable compounds common to both horses and humans [29-31]. Metabolic consistency was strongest across the plasma profiles. This validates the concept of 267 268 conservation of mammalian physiology across species and that homeostatic metabolic control is tight in both systems. A lower percentage of the equine faecal and urine metabolic profiles 269 270 were shared with humans but this was to be expected due to different digestive systems and 271 metabolic pathways. In contrast, ruminants share a number of gut microbial co-metabolites 272 such as hippurate and PAG [32-34]. These herbivores have similar digestive strategies to 273 obtain nutrients from a cellulose-rich diet. 274 Excretory biofluids (urine and faeces) are the most metabolite-rich. The environmentallydetermined nature of these biofluids renders them under less tight homeostatic control than 275 276 that of the plasma. However, urine was also remarkably tightly controlled, given the 277 relatively homogeneous genetic and environmental backgrounds of Thoroughbred racehorses. 278 Differing creatinine excretion in one animal (due to the sample being collected after exercise) was responsible for the increased inter-animal variation observed (Fig 4A). In urine, 39 279 280 metabolites were identified as being unique whereas 21 metabolites were faeces-specific and 281 12 metabolites were plasma-specific. However, there were a small number of low-282 concentration unassigned metabolites that have not been included in these counts. All 3 283 biofluids are likely to contain thousands of metabolites which are too dilute to produce a 284 significant NMR signal or were not detected by NMR in this study, but we feel we were able

to capture a good representation of the normal equine metabonome and this has been verified by our ongoing work on a larger cohort of animals (data not shown).

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Inter-compartmental variation is greater than inter-animal variation and is consistent with other mammalian studies [8; 35]. Faecal samples varied from other biological matrices due to the presence of a number of SCFAs, formate and isovalerate. SCFAs are the product of gutmicrobial fermentation of dietary fibre and contribute up to 70% of a horse's energy requirements [36; 37]. Acetate level variability is likely to be due to differing bacterial communities and consequently SCFA production. Collection time in relation to feeding time is known to exert an effect on SCFA levels in equine faeces [38]. Samples were consistently taken before a morning feed. However, this variability could be due to the difference in individual intestinal transit times [39; 40]. Other SCFAs were not seen to vary to the same extent as acetate. Butyrate is the main energy source for colonocytes and hence is likely to have been utilised rather than excreted [41]. Butyrate and propionate are extensively metabolised by first-pass metabolism and therefore absent in NMR detectable quantities in urine and plasma samples. Creatinine was the greatest source of variation amongst urine samples. Creatinine is a waste product of muscle metabolism formed from creatine in order to maintain ATP levels during exercise [42]. The concentration of urine and thus metabolites can change dramatically. Normalisation of the data prior to analysis helps to minimise spectral anomalies caused by differences in urinary dilution. Creatinine levels can vary according to factors such as muscle mass, physical exercise, diet, age and muscle damage from previous strenuous exercise [43].

Urine was found to differ from the other biological matrices due to the presence of a number of gut-microbial co-metabolites (hippurate, PAG and *p*-cresyl sulphate and TMAO). These

The urine metabolome provides a metabolic window into gut microbial co-metabolism.

compounds originate from exogenous sources (microbial and dietary) and are incorporated
into the host circulation after absorption. They subsequently undergo enzymatic conjugation
in the liver and gut mucosa to increase their polarity and enable renal excretion. In mammals,
enterohepatic recycling means further metabolism can occur at the gut-level [44]. Hippurate
is an aromatic compound predominantly formed from glycine conjugation of dietary or
microbial benzoate and PAG is a glycine-conjugated microbial metabolite of phenylalanine
metabolism (glutamine in higher apes and humans) [45]. P-cresol is formed from bacterial
degradation of tyrosine and is subsequently sulfated or glucuronidated in the liver or gut [46].
Both forms of conversion occur in rodents and horses, whereas humans predominately form
sulphate conjugates and pigs predominately glucuronidate the cresol molecule.
Trimethylamine (TMA) is a microbial degradation product of dietary choline; this metabolite
is absorbed from the gut and subsequently oxidised in the liver to produce TMAO [47]. A
number of anaerobic bacterial populations are known to produce these metabolites including
clostridia [48-50]. This taxonomic classification of bacteria has been associated with
intestinal disease in horses as well as being part of the normal microflora [51-54].
¹ H NMR spectroscopy is a robust method for assessing the inter-animal variation in
Thoroughbred racehorses. It is important to address potential sources of variation when
assessing metabonomic studies. The sensitive nature of such investigations necessitates the
collection of metadata to help explain possible variation between the samples. This
information should include details from sample collection, sample storage and run order, as
well as information relating to sample subject (health status, age, sex for example). The effect
of these is widely reported in the literature [35; 55; 56]. Although the 7 horses were taken
from 2 different yards, the samples clustered tightly, highlighting the metabolic uniformity of
these Thoroughbred racehorses despite differing age ranges, location and stages of fitness

(see Supplementary Item 1). This study was intended to create a reference tool for research into Thoroughbred racehorse metabolism and microbial co-metabolism. Since the samples from this first pilot clustered closely for all 3 biofluids indicating that the dominant metabolites visible by NMR were conserved across animals, it was deemed unnecessary to collect samples from further animals to minimise unnecessary sampling. Although, the sample size in this study is small and only one metabolic snap shot was taken for each horse, the fact that inter-animal variation in metabolic profiles was low, suggests that 'healthy' Thoroughbreds share a similar metabolic phenotype, which we explored using a range of NMR-based structural elucidation tools including 2-D pulse sequences to elicit carbon-proton correlations and statistical spectroscopy methods. Our future work includes a larger study investigating normal variation amongst different racehorse populations over time. This work comprehensively assigns dominant features of the ¹H NMR spectra of the equine metabonome from plasma, urine and faeces and for the first time provides baseline information for future studies in equine health and disease. Urine and faecal profiles provide an insight into host-microbial metabolic interactions, whereas plasma profiles are more likely to represent host physiological processes. The purpose of this study is to provide an analytical template to researchers thinking about adding metabonomic analysis to their experiments and to indicate which biofluids may be of use to them. We have showed that faecal samples are more variable that either urine or plasma. Plasma is under tight homeostatic control and thus might be expected to show relatively less variation. However, urine, other than one identified outlier, exhibited less variation than the other biofluids under investigation and contributed the greatest number of identifiable metabolites. Further studies are warranted using metabonomic and metagenomic technology to explore the role of gut microbes on equine physiology and metabolism.

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359	Authors' declaration of interests
360	No competing interests have been declared.
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366	Sampling and metadata collection were carried out under University of Liverpool ethics
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368	
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373 374	
375	Authorship
376	E. Escalona contributed to all sections. J. Leng contributed to study execution, and data
377	analysis and interpretation. J. Swann and A. Dona contributed to study design, study
378	execution, and data analysis and interpretation. C. Merrifield contributed to the preparation of
379	the manuscript. All authors gave their final approval of the manuscript.
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384 Figure legends:

Fig 1: (A) 700 MHz 1D ¹H-NMR spectrum of urine, (B) CPMG ¹H-NMR spectrum of plasma, (C)1D ¹H-NMR spectrum of faecal water. All spectra partially labelled according to the assignments made in Table 1 and Supplementary Item 2.

Fig 2: (A) 800 MHz ¹H COSY NMR spectrum of the aromatic region of urine (B) J-res NMR spectrum of plasma highlighting the aliphatic region between 3-4.5 ppm. Key to metabolite identification is provided in Table 1 and Supplementary Item 2.

Fig 3: (A) Venn diagram highlighting the degree of metabolite overlap between biological compartments. The central section represents the number of core metabolites visible across all biological matrices (14 ubiquitous metabolites). The outer circle with numbered slices represents biological matrix specific metabolites that are ordered according to origins/function, and numbered according to metabolites in Table 1. Compartments are colour coded - plasma (red), urine (yellow) and faeces (green). Coloured dots represent metabolic functions, protein and amino-acid metabolism (purple), energy metabolism (orange) and fat metabolism (blue). Asterisk denotes metabolite can be mammalian in origin and a 'd' denotes metabolite can be from dietary origin.

Fig 4: (**A**) PCA scores plot demonstrating increased faecal variability relative to other biological matrices. Plasma (red), urine (yellow) and faecal (green) samples. Principal component 1 (PC1) accounts for 51%, PC2 for 15%, and PC3 for 5% of the total variation. Variance colour plots indicating the percentage of the total variance for each metabolite in (B) faeces and (C) urine (Mean spectrum is plotted, coloured by variance expressed as a percentage of the total variance).

Table 1: Metabolites identified using ¹H-NMR of equine plasma, urine and faeces. 410 Metabolite numbers correspond with annotated ¹H-NMR spectra in Figs 1, 2 and 3. 411 Metabolites are assigned to biofluids denoted by coloured dots; urine (yellow), plasma (red), 412 413 faeces (green). The 14 core metabolites are highlighted in purple. 414 415 416 Manufacturers' addresses 417 ^aBertin Technologies, Montigny-le-Bretonneux, France 418 ^bBruker, Massachusetts, USA 419 ^cMATLAB, Mathwork, Massachusetts, USA 420 421 Additional Supplementary Items may be found in the online version of this article at the 422 publisher's website: 423 424 **Supplementary Item 1:** The table highlights sample metadata and possible sources of 425 variation within the data set. Yard, gender, age and training schedule are included. Gender is 426 denoted as G=gelding, C=colt. 427 Supplementary Item 2: Table showing metabolites found in plasma (red circle), urine 428 (yellow circle) and faeces (green circle) using ¹H-NMR spectroscopy. Peak multiplicities and 429 chemical shifts are shown and structural information is also provided. † indicates tentative 430 431 assignment. Details on each metabolite's origin and function are highlighted as well as a link 432 to the metabolite's page in the hmdb database. 433 434 **Supplementary Item 3:** Table denoting feeding regimes on the 2 yards sampled. Top section 435 highlights concentrate feeds and bottom highlights roughage types offered. 436 **Supplementary Item 4:** 800 MHz ¹H COSY NMR spectrum of the aliphatic region of 437 438 faeces, between 3-4 ppm. Key to metabolite identification is provided in Table 1 and 439 Supplementary Item 2. 440

- **Supplementary Item 5:** 800 MHz ¹H ¹³C HSCQ NMR spectrum of the aromatic region of
- 442 urine, between 8-7 ppm. Key to metabolite identification is provided in Table 1 and
- Supplementary Item 2.

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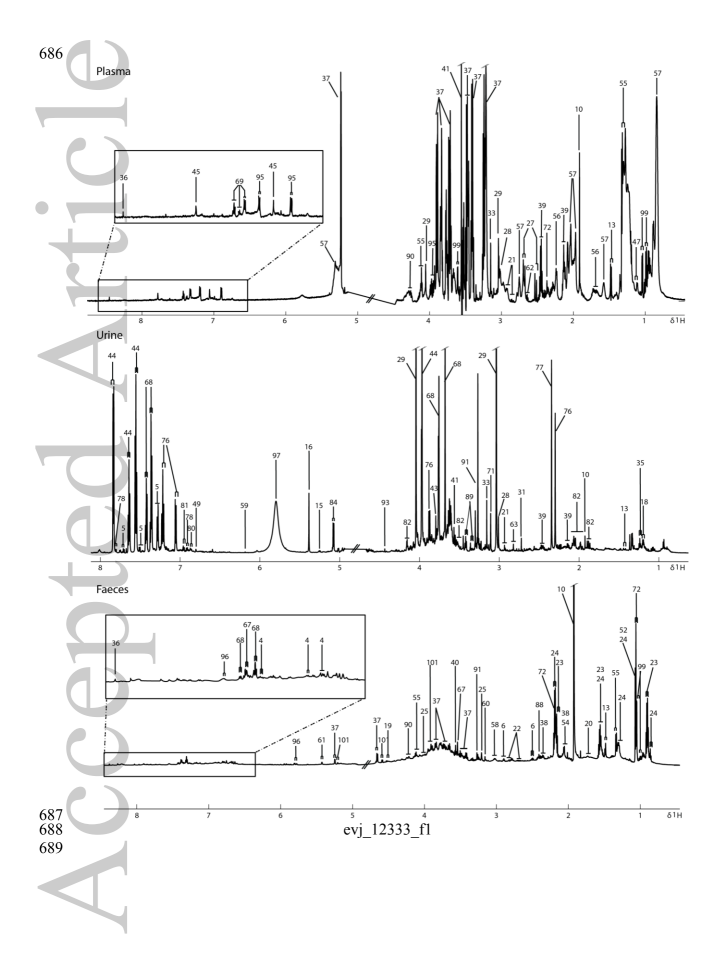
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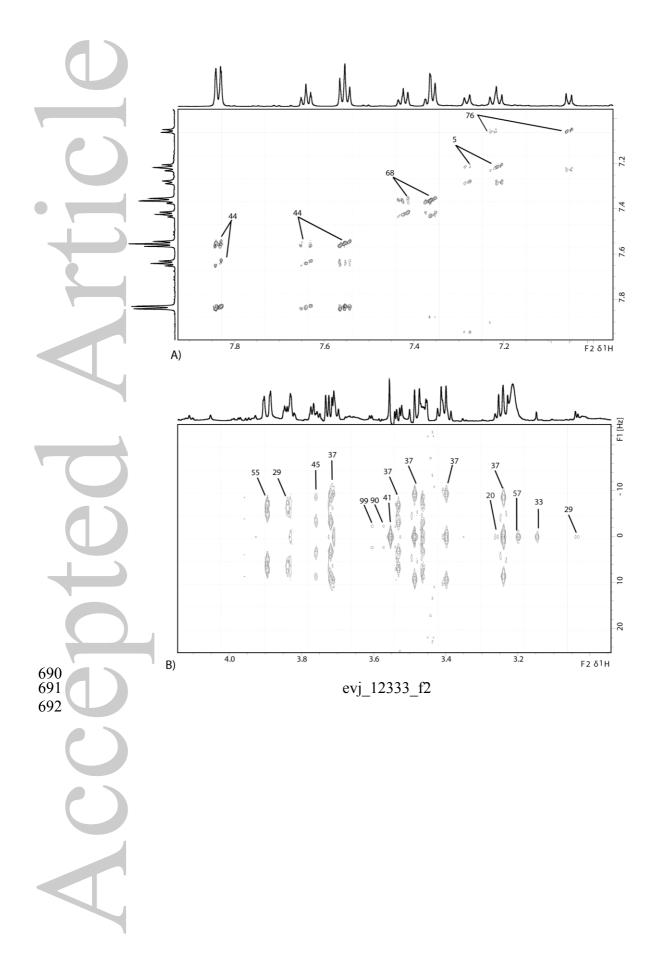
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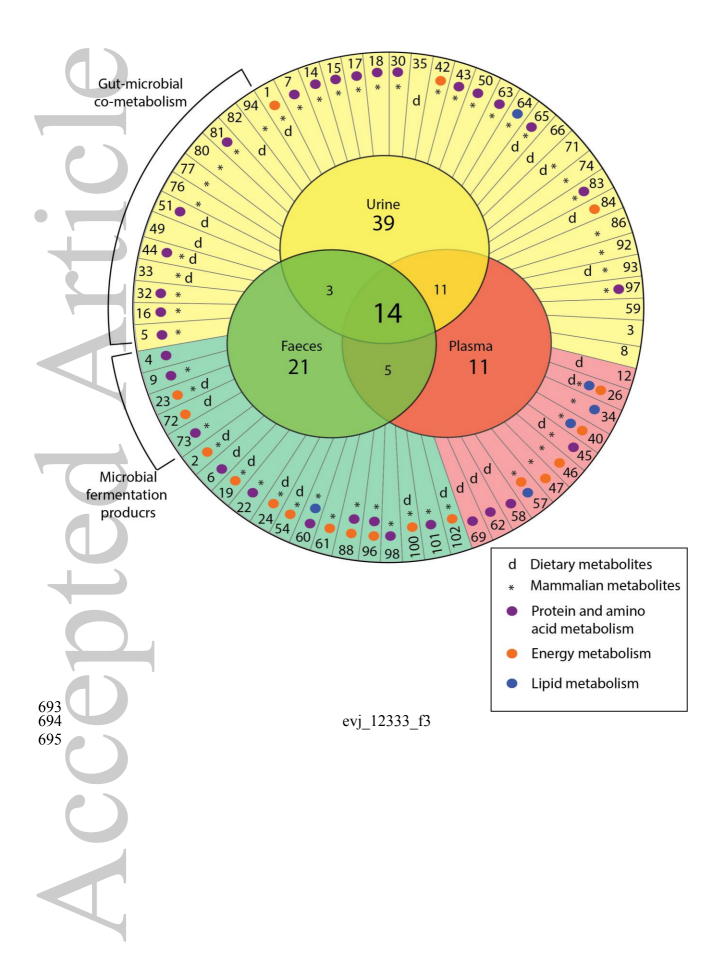
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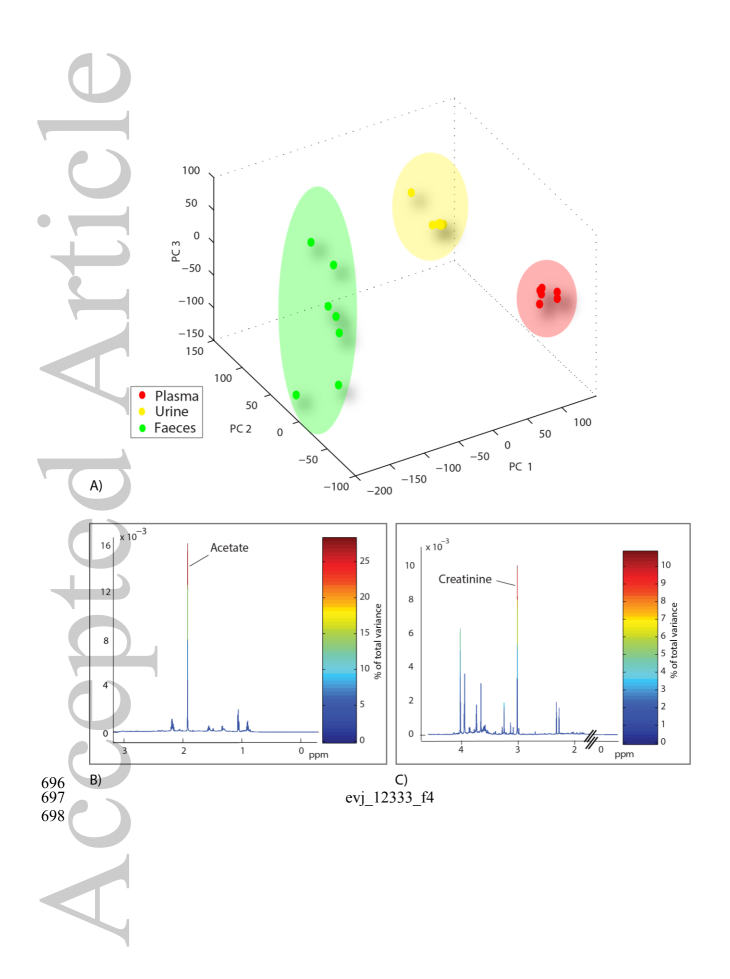
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N°	Compound
1 🧶	2-Hydroxy-3-methylbutyric acid
2	2-Methylbutyrate
3 🔾	3-hydroxy-4-methoxymandelic acid
4 🔍	3-Hydroxyphenylacetate
5 🔘	3-Indoxyl sulphate
6	3-phenylpropionoate
7 🔍	3-Ureidopropionic acid
8	4-hydroxy-3-methoxymandelic acid
9	5-Aminovalerate
10	Acetate
11	Acetoacetate
13	Adipate
13 🛡 🔍	Alanine
14	β Alanine
15	Allantoate
16	Allantoin
17	α-Aminoadipate
18	Beta-Aminoisobutyrate
19 🔍	Arabinose
20 0 0 21 0 22 0 23 0	Arginine
21 🛡 🔾	Asparagine
22	Aspartate
23	Butyrate
24	Caprylate
25 🔎 🔍	Choline
26	Cholestrol
27 🔍 🔾	Citrate
28	Creatine
29 🛡 🔾	Creatinine
30 31	Dihydrothymine
31	Dimethylamine (DMA)
32	Dimethylglycine (DMG)
33	Dimethyl sulphone
34	Ethanolamine
35	Ethyl glucoside
36	Formate
37	Glucose
38	Glutamate
39	Glutamine
40	Glycerol
41	Glycine
42	Glycogen
43	Guanidoacetate
44 0	
44	Hippurate Histidines
45	
46	α Hydroxybutyrate
47	β Hydroxybutyrate
48	Alpha- Hydroxyisobutyrate
49	Hydroquinone
50	Hypotaurine
51	Indole-3-acetate

N°	Compound
52	Isobutyrate
53 • •	Isoleucine
	Isovalerate
55	Lactate
56 •	Leucine
57	Lipids
58	Lysine
59	Maleic anhydride
60	Malonate
61	Maltose
62	Methionine
63	Methylguanidine
64 0	Nicotinurate
65	Orotate
66	Pantothenate
67	Phenylacetate
68	Phenylacetylglycine (PAG)
67	Phenylalanine
69	Proline betaine
70 •	Phosphocholine
71	Proline betaine
72	
73	Propionate
74 0	Putrescine
75 • •	Pyridoxine
	Pyruvate
76 🔍	p-cresol glucuronide
77 🔍	p-cresol sulphate
78	<i>p</i> -hydroxybenzoate
79	<i>p</i> -hydroxyphenylacetate
80	<i>p</i> -hydroxyphenyllactate
81 🔍	<i>p</i> -hydroxyphenylpyruvate
82 🔾	Quinate
83 🔍	Quinone
84 🔍	Raffinose
85 🔍	Scyllo-inositol
86 🔍	Sebacate
87 🔍	Serine
88	Succinate
89	Taurine
90 0	Threonine
91 • •	Trimethylamine-N-oxide (TMAO)
92 🔍	Trans-aconitate
93 🔵	Trigonelline
94 🔍	Trimethylamine (TMA)
95 🔵 🔘	Tyrosine
96 🔍	Uracil
97 🔍	Urea
98	Uridine
99 🔵 🔾 🔘	Valine
100	Valerate
101	Xanthine
102	Xylose
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