From Saliva to Faeces and Everything in Between: A Guide to Biochemical Analysis Using Animal Samples for Biomarker Detection

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Over the last decade, interest in the emotional states and stress levels of animals has grown. These emotional states can have secondary effects for owners; for example, if an animal becomes aggressive, this can lead to relinquishment or even euthanasia. In addition, long-term stressful situations can have serious health impacts on animals and can affect meat quality in livestock. A variety of methods can be used to investigate biomarkers in animals, and many sample types can be taken to facilitate this. The choice of assay will often depend on the animal under investigation and practicalities of obtaining the sample. The assay choice can also be dependent on testing conditions such as the field versus laboratory, samples taken, costs, and the desired results. There is also the question of the timescales of the investigated response: Do you want to test what happened over the last month? Last week? Yesterday? Or within the last hour? This review highlights some of the pros and cons of the different samples and the different methods for biomarker analysis in animals. Studies can be made or broken based on the type of samples taken, and what aspects are to be investigated, and this simple decision can make a world of difference to the results of an investigation. Careful planning and thought before starting a study can make the difference between a scientific breakthrough with animal welfare and husbandry implications and poor results that are of little use to man or beast.

Keywords: biochemical analysis, ELISA, saliva, blood, hair, faecal

Introduction

Careful selection and analysis of biological samples is vital for accurate quantification of animal health, condition, and welfare. This review covers some of the key considerations for selecting the biological sample for analysis and the type of biochemical test(s) required to analyse the biological markers (biomarkers) present. As an extra complexity, multiple biomarkers can be used for similar types of analysis; for example, to evaluate stress, cortisol, amylase, or even IgA are potential biomarkers (Hong et al., 2019; Mack & Fokidis, 2017; Muneta et al., 2010). As you would imagine, these different biomarkers are all present in the host animal, but they will be at different levels or even totally absent depending on the choice of biological samples (blood plasma, saliva, urine, faeces, etc.). The type of biological assay, or the number of steps needed to process/extract the biomarker, varies both with the type of sample and with the selected biomarker. These biomarkers are often hormones but can also be enzymes, other proteins, or metabolites (including hormone metabolites). These factors might even require compromises between selecting accuracy over easy, rapid, or in-field testing, when selecting the sample and/or biomarker to be investigated.

This review is divided into two major sections: The first section covers the aspects of selecting samples, different sample types, and examples of biomarkers that can be measured from these samples. The second section mostly focuses on the different types of biochemical assays that can be used to measure biomarkers and simple descriptions of how these assay work or may be used. Given the
enormous breadth of the review, we have provided general examples and citations for more specific examples covering the major sample types, commonly used biomarkers, and biochemical assays. The goal of the review is to provide a first resource for researchers interested in studying biomarkers to complement behavioural analysis. This article should not be considered a complete guide. We urge readers who are planning to use any of the techniques described here to read more detailed information in the many publications we have cited, because many of our general rules of thumb may not apply to specific implementations.

**Choosing the Ideal Sample**

Selecting the ideal physical sample is incredibly important. Although most assays are available to run either on site or in a laboratory, this may not be the case for all samples. The selection of sample type is dependent on the biomarker(s) to be measured, the types of assays available to measure these biomarkers (see the Types of Biochemical Test section for further information), and the availability to collect the sample type. Availability will depend on many factors, not least the ability to collect invasive samples, such as blood, and the research project questions/aims (Mormède et al., 2007). It is important to mention that the correct ethical and legal licences must be in place before samples are obtained from animals. In addition, licences may also be needed for working with certain species, such as Convention on International Trade in Endangered Species of Wild Fauna and Flora (CITES) species, or for transportation of samples across borders or into countries because of biosecurity concerns (Animal Science Procedure Act [ASPA], 1986). As an example, a schematic diagram for selecting the ideal sample for studying stress responses is shown in Figure 1.

**Blood**

Blood is generally regarded as the gold standard sample, but sampling is highly invasive and as such may require licenses, such as a Home Office Licence in the UK (ASPA, 1986). It is also important to remember that the process of taking blood can alter the level of stress hormones in the animal, particularly when an animal is not keen on being handled (e.g., sheep; Romero & Read, 2005). In other cases, when animals require capture in the wild, the stress of the process may induce glucocorticoid hormone release, so ideally samples should be taken within a few minutes of capture while ensuring sample collector safety (Romero & Read, 2005).

Blood samples have to be processed correctly—commonly by low-speed centrifugation or clotting into sedimentation layers such as the cells, buffy coat, or plasma/serum—to ensure that the layer with the biomarker is retained (Bielohuby et al., 2012). It is also worth noting that a substantial difference exists, especially for protein concentrations such as fibrinogen, between serum and plasma, so care must be taken to ensure that the correct one of these two is chosen (plasma has fibrinogen and clotting factors, but they are absent or greatly reduced in serum samples).

Blood samples can be stored with relative ease at room temperature or chilled while sampling, before ideal storage at −80 °C for sera and plasma (Bielohuby et al., 2012; Lombardi et al., 2012). However, biomarker stability can vary with temperature, allowing conformational changes or proteolytic cleavage (Reimers et al., 1983). Freeze–thaw cycles can also cause degradation of some hormones, so ideally, the samples that are to be used to detect multiple hormones should be aliquoted out into several smaller volumes for single use (Bielohuby et al., 2012).

It is also important that dilution of blood or serum sample, even by introduction of an anticoagulant as commonly occurs in a vacutainer, is taken into account with final calculations, as it may affect the results (Bowen et al., 2010; Kontrn et al., 2011; Lippi, Franchini, et al., 2006). Haemolysis, which can be caused by inappropriate sample handling and storage, can also lead to dilution and potential contamination of the sample and should be prevented where possible (Bellomo et al., 2012; Koseoglu et al., 2011; Lippi, Salvagno, et al., 2006).

Additionally, and importantly, there is a difference in measurable blood analytes, especially with rodents, for which there is a large difference in the site from which the blood sample is obtained (e.g., tail vain, tail tip sampling, jugular vein, cardiac puncture; Aasland et al., 2010; Arola et al., 1980; Christensen et al., 2009; Fitzner Toft et al., 2006; Vahl et al., 2005). Therefore, it is worth considering where you would take the terminal blood sample from if an animal is to be euthanised.
There are also biomarker variations on a daily, monthly, postfeeding, and even sometimes yearly circadian rhythm with hormones such as testosterone and progesterone, as has been shown in maned wolves, and in seals (Greig et al., 2007), which show increased peaks in certain seasons (Maia et al., 2008).

Although blood is a complicated mixture of a large number of proteins, it offers a rapid or even instantaneous insight into stress hormones. Indeed, it may be so fast that unless samples are obtained very rapidly, the stress of being handled may create issues and inflate stress hormone levels.

**Saliva**

Saliva is a relatively easy-to-obtain sample and can contain various hormones, including cortisol (Cobb et al., 2016; Cook et al., 2013; Wenger-Riggenbach et al., 2010). As cortisol is a lipid soluble hormone, it crosses cellular membranes, allowing for its detection in saliva within 15 min of a stressor being applied, making it an ideal sample for rapid stress hormone detection (Dickerson & Kemeny, 2004; Gunnar & Vazquez, 2006; Kirschbaum et al., 1993; Shirtliff et al., 2015). This rapid release and ease of testing has led to development of point-of-care devices to assess stress in both humans and animals (Choi et al., 2014; Kaushik et al., 2014; Nara et al., 2010; Zangheri et al., 2015).

Depending on how sampling is performed, it may also require a licence. For example, within the United Kingdom, the ASPA (1986) means that you can swab around the outside of the teeth but a licence is required to enter the buccal cavity, as this is considered an invasive sample. In addition, the volume of saliva can sometimes be small, from around 100 µl from dogs (personal experience) to around 500–750 µl from equids using the EquiSal® saliva collection swab (Austin Davis Biologics Ltd., Northamptonshire, UK).

A wide range of studies have used saliva from various animals to test for different hormones associated with behaviour and stress, including cortisol (Cobb et al., 2016; Cook et al., 2013; Wenger-Riggenbach et al., 2010), luteinizing hormone (Srinivasan et al., 2020), oxytocin (MacLean et al., 2018), vasopressin (MacLean et al., 2018), prolactin (Gutiérrez et al., 2019), and testosterone (Kutsukake et al., 2009).

Studies have also shown a strong correlation between cortisol in blood and in saliva, suggesting that saliva may be a useful proxy for blood sampling, avoiding invasive sampling (Fell et al., 1985; Greenwood & Shutt, 1992; Neigrão et al., 2004), although other studies disagree (Dzviti et al., 2019). It does have the advantage of being repeatable multiple times in a relatively short period (Koyama et al., 2003). However, this can be difficult in field situations for wild animals, but novel methods of saliva collection have been devised and used successfully (Higham et al., 2010; Smiley et al., 2010).

Care also needs to be taken when choosing the method for saliva collection. Potentially, swabs can be contaminated with food debris, and even plant hormones can interact with immunoassay antibodies (Dabbs, 1991; Granger et al., 1999), and some biomarkers may adhere to the cot-

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**Figure 1.** A flowchart diagram to aid with the selection of the ideal sample for the study which you wish to undertake. Note that this may differ from country to country as licencing requirements vary. Y = yes; N = no.
ton swab and thus escape detection (Shirtcliff et al., 2001). Processing of saliva can be difficult if the sample is highly viscous, and samples can precipitate after a freeze–thaw cycle (Read et al., 1990; Riad-Fahmy et al., 1982). Once again, several freeze–thaw cycles can alter the concentration of hormones (Gröschl et al., 2001).

Additionally, salivary flow rate can influence results by diluting potential hormones of interest. This can be exacerbated by many factors, including increased temperature, food and liquid intake (Elmér & Ohlin, 1971; Ito et al., 2001; Jacques et al., 1989), and exercise (Colussi et al., 2018). Food intake can cause errors in hormone measurement (Laudenslager et al., 2006; Magnano et al., 1989), and care must be taken to avoid causing bleeding and getting blood contamination of the swab, as this can alter the hormone concentration (Dzviti et al., 2019). Even an easy-to-obtain sample may require considerably more planning than one might think (Kalliokoski et al., 2019; Koren et al., 2019).

### Hair or Other Tissue

Hair and tissue are rapidly developing areas of interest for studies associated with hormones and behaviour/cognition. Hair presents a simple, relatively easy-to-obtain sample but can prove difficult if working with certain animals, such as wild cats, because of concerns about collecting samples safely.

Although mainly used for human medicine so far, hair can act as a nice proxy for long-term analysis of hormones that are encompassed into the hair during its growth (Schweikert & Wilson, 1974). Studies often focus on a variety of hormones, including testosterone, oestriadiol, cortisol, progesterone, cortisol, and androstenedione (Chen et al., 2013; Gao et al., 2013, 2016; Grass et al., 2016; Kapoor et al., 2014; Yang et al., 1998).

Complications exist with using hair for analysis of some hormones; for example, cortisol levels can vary with the hair’s location on the body, so standardisation of sampling area is important to allow for comparisons (Carlitz et al., 2015; Heimbürge et al., 2019; Mesarcova et al., 2017; Terwissen et al., 2013; Yamanashi et al., 2013). Thus, collection of random hair from nests, traps, or fences, such as from rodents or sheep, can prove problematic (Heimbürge et al., 2019). The reasons for these variations remain unclear but may be associated with factors such as the level of sunlight to which a body part is exposed (Grass et al., 2016).

Hormone degradation does not appear to be a great concern with hair, meaning that the sample can easily be stored at room temperature without the need for desiccation or freezing (Accorsi et al., 2008), but washing hair samples is important to limit contamination with sweat, urine, or faecal matter (Chen et al., 2013; Ferrero & Liberles, 2010; Gao et al., 2016; Musshoff & Madea, 2007; Sheriff et al., 2011).

Hormone concentration can vary with factors that influence the growth rate of the hair; these factors can include age, part of the body, sex, breed, and species (Heimbürge et al., 2019). In addition, concentrations can vary with hair colour, as seen in dogs and chimpanzees (Bennett & Hayssen, 2010; Taylor et al., 2015; Wennig, 2000; Yamanashi et al., 2013). Questions also remain about the effectiveness of measuring hormones in hair strands. Most hormones are concentrated in the bulb of the hair and not completely incorporated into the hair strand as it grows, so hormone concentration is highly dependent on the part of the hair that is analysed (Keckeis et al., 2012; Stubsjøen et al., 2015).

As an alternative to hair, a sample of claw tissue, such as that obtained during routine trimming of nails in animals, may be used. This tissue provides assessment of stress hormones in the animal over a long period (Matas et al., 2016). A correlation has been shown between hair and claw cortisol in newborn dogs (Veronesi et al., 2015). For avian species, feathers can be used rather than hair (Matas et al., 2016).

Thus, hair, feathers, and claws can offer a simple, and in some cases easy-to-collect, sample, but some level of planning is required to ensure that results obtained from these samples are comparable and that differences are not due to different sample types.

### Faeces

Faeces is possibly the easiest sample to obtain, as it is left naturally by all animals. Perhaps an obvious statement, faeces is a dirty sample filled with various hormones and enzymes at different stages of metabolism, animal gut microbiota, food remains, and other bits that may interfere or inhibit certain reactions such as polymerase chain reaction (PCR; El-Bahr et al., 2005; Lepschy et al., 2007; Palme, 2005). Indeed, differences in the gut microbiota can lead to differences in hormone metabolism and breakdown, so this may affect results of faecal hormone analysis (Antwis et al., 2019; Hooda et al., 2013).

Faecal sampling has proven particularly useful for different research focuses, including long-term stress, seasonal hormone patterns, and pregnancy status (Cizauskas et al., 2015; Garnier et al., 1998; Hadinger et al., 2015; Hernandez et al., 2018; Isobe et al., 2005; Nemeth et al., 2016, Schwartzzenberger et al., 1996; Wasser et al., 1991; Wheeler et al., 2013). This sampling is commonly used...
to investigate the presence of steroidal hormones, such as oestrogens, androgens, and progestins for reproductive status determination and glucocorticoids for stress analysis (Amaral, 2010; Goymann et al., 2002; Harper & Aus- tad, 2000; Kotrschal et al., 1998).

However, the use of faeces for hormone detection is somewhat limited by heavily degraded samples because of the harsh conditions of the gut as well as the actions of the bacteria in the microbiome (Antwis et al., 2019; Hooda et al., 2013). As such, it is important that faecal samples are collected rapidly postdefecation and frozen to prevent further bacterial degradation of hormones (Hodges & Heis- termann, 2011; Millspaugh & Washburn, 2004), as this will affect the results (Palme, 2005, 2019). Where freezing is not possible, such as in field situations, rapid drying the sample may prevent significant degradation, such as can be achieved by using alcohol to remove water (Palme, 2005), but this is not always ideal. Addition of sodium azide or other acids can aid preservation of hormones in alcohol at room temperature, although these harmful chemicals are dangerous to the environment and may not be suitable in the field (Whitten et al., 1998). More recently, rapid field extraction has given a more accurate picture of the hormones that are present in a faecal sample (Beehner & Whitten, 2004; Kalbitzer & Heistermann, 2013; Whit- ten et al., 1998; Ziegler et al., 2005).

Biomarkers are not always evenly distributed throughout a faecal sample, so the part of the sample that is taken needs to be carefully planned. This is particular- ly true for large animals, which produce a large faecal sample, such as cattle and elephants. The same part of the sample should be taken to allow for appropriate comparison between samples (Hadinger et al., 2015; Peter et al., 2018). Then homogenisation is required prior to analysis; removal of large pieces of nondigested foods such as seeds and insects will aid in this (Millspaugh & Washburn, 2004; Palme, 2005).

Difficulties can also arise in collection and potential contamination, especially urine contamination, which can increase concentrations of some hormones, skewing results (Hay et al., 2016; Schönnig et al., 2002). Despite difficulties, studies have been carried out on aquatic animals (Am- aral, 2010). A major positive to this sample is that there are limited diurnal variations of hormones within faeces, unless there is a very rapid gastrointestinal tract transit time (Goymann, 2005, 2012; Millspaugh & Washburn, 2004).

Although a simple sample to collect in many cases, faeces needs some thought and some standardising to ensure that the results can be appropriately analysed and compared.

**Urine**

Urine is a mixture of waste products and hormones that are commonly excreted as conjugated water-soluble forms. They are often an indication of the presence of hormones over a few hours rather than an instantaneous indication as seen with saliva (Bouatra et al., 2013).

Urine is already widely used for detection of pregnancy hormones in humans, and the detection of human chorionic growth hormone (Chard, 1992) is used in home pregnancy test kits. Urine is also useful in disease diagnostics, such as monitoring diabetes in humans and in dogs (Hess et al., 2000).

Urine can be relatively easy to obtain from some animals (e.g., through a swab of the bottom of the cage) but relatively difficult from others and may require some training of the animals (Kurien et al., 2004; Laule et al., 1996). When working with wild animals, collection of this sample can be even more difficult, but methods have been developed (Daniéh et al., 2015; Knott, 1997). Urine has been used to detect a variety of peptide hormones, such as cau- xin (McLean et al., 2007), trypsinogen activation peptide (Allen et al., 2006), e-peptide (Polonsky et al., 1984), oxy- tocin (Mitsui et al., 2011; Nagasawa et al., 2009), vasopressin (Dolph et al., 1962), and thyroid hormones (Goff et al., 1986), as well as certain immune parameters, particularly linked with infection with various pathogens (Dor- ward et al., 1991; Ravnik et al., 2014).

Although some biomarkers are considered to be stable in urine at room temperature for up to 24 hr, some degradation may occur because of the bacteria within the sample (Grant & Beastall, 1983). Equally, urine samples should be centrifuged to remove particulate matter, and care should be taken when freezing urine samples, as this may lead to further sample degradation (Heistermann, 2010).

One should bear in mind that the volume of urine, which is linked to the hydration of the animal, will have an impact on the concentration of hormones. For example, a well-hydrated animal will produce higher volumes of urine than a dehydrated animal, and more di- lute urinary hormones (Miller et al., 2004). To counter this issue, creatinine is sometimes used as a counter- balance, but this is also subject to many fluctuations based on diet, age, sex, breed, kidney function, and so on (Carrié et al., 2000; Miller et al., 2004; Thompson et al., 2012).

Urine also allows for an investigation into varying concentrations of hormones over a day, with a sample that is easier to collect than saliva in some cases (Heis-
terminann, 2010). However, this can cause issues, as certain hormones can be naturally increased at certain times of day. This increase may not be in response to a stressor; for example, in some species of monkey cortisol levels can be higher in the morning than at other times of day (Davis et al., 2005; Muller & Lipson, 2003; Smith & French, 1997). This is also true for dogs, which show variation during the day and night (Gordon et al., 1985; Kolevská et al., 2003). Therefore, throughout the study the timing of sampling should be as close to the same as possible. If similar timing cannot be maintained, urinary hormones should be analysed with caution.

Although a urine sample is easy to obtain for some animals, several other aspects need to be factored into studies to make sure that the results are reliable and easy to interpret. Whatever sample is chosen, the extraction method is important. Method is discussed later in this article.

Types of Biochemical Tests and Their Applications

Commonly, the selection of a biochemical test is determined by the choice of biomarker and the type of sample available for its isolation. Key considerations in selecting the most appropriate test(s) are briefly discussed next and summarised in Table 1. These considerations include the type of biomarker (chemical, protein, DNA/RNA), how much sample can be collected (as some methods require as little as 1–10 µl of sample but other tests require several milliliters), budget, and whether the test needs to be performed in situ.

<table>
<thead>
<tr>
<th>Test Type</th>
<th>Biomarker</th>
<th>Method Output</th>
<th>Limitations</th>
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<tr>
<td>Chemical</td>
<td>Chemicals and proteins</td>
<td>Mostly colorimetric reporter systems or colour change</td>
<td>Sensitivity of the test. Amount of sample required.</td>
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<tr>
<td>Enzyme-linked assay</td>
<td>Chemicals, proteins</td>
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<td>Sensitivity of the test. Amount of sample required.</td>
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<tr>
<td>Enzyme activity</td>
<td>Enzymes</td>
<td>Mostly colorimetric reporter systems</td>
<td>Sensitivity of the test. Amount of sample required.</td>
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<tr>
<td>PCR test</td>
<td>RNA or DNA</td>
<td>Graphic or gel band. Indicator of presence or copy number.</td>
<td>Availability of RT-PCR facilities/equipment. Ability to extract RNA/DNA from the samples.</td>
</tr>
<tr>
<td>Proteomics</td>
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<td>Mostly colorimetric reporter systems or identification of modifications by affinity isolation then either mass spectrometry or western analysis using PTM specific antibodies</td>
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</tr>
<tr>
<td>Mass spectrometry</td>
<td>Chemicals and proteins</td>
<td>Graphic/tabulated data. Indicator of presence or amount present.</td>
<td>Potentially required as an external service because of expertise and equipment cost</td>
</tr>
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Chemical Assays

Chemical detection of biomarkers directly measures a chemical in a biological sample by causing a reaction to report the presence or amount of the chemical. Chemical assays are routinely used to detect and measure the levels of sugars or proteins in urine, for example (Boag et al., 2019; Shropshire et al., 2018). They are comparatively inexpensive compared with antibody test assays, which are quick to perform but can lack sensitivity. An example of rapid, easy-to-use chemical assays are “dipstick” assays. A “dipstick” is dipped briefly (usually for 1–15 s) into the biological liquid being assayed, then left for about 15 s for the reaction to occur. The readout is colorimetric, and the amount of colour produced is proportional to the amount of chemical present. The systems used for diabetes testing in animals is a common example. The dipstick colour can be compared with a colour range chart supplied with the assay kit to provide semiquantitative data, which allows for rapid in-field measurements of proteins, chemicals, and pH by comparatively untrained researchers (Athanasiou et al., 2018; Boag et al., 2019).

Enzyme-Linked Assays

Enzyme-linked assays are commonly used either as a reporter system “linked” to another chemical reaction, in order to get a quantitative readout, or to amplify the readout of another reaction to increase assay sensitivity. Enzyme-linked assays are, in general, more complex than standard chemical or enzymatic assays. They are often used to measure enzymatic activity and have the ability to amplify the reporter system so that low levels of activity can be more easily measured. These assays have
multiple stages and require dedicated laboratory systems/equipment. Not all enzyme-linked assays require the use of antibodies but simply use linked enzyme systems as a method of reporter amplification for detecting the activity of enzymes at very low concentrations. Antibody detection systems such as Enzyme Linked Immunosorbent Assays (ELISAs) often rely on enzyme-linked reporter systems to amplify the colorimetric quantitative readout systems. These ELISA systems are becoming veterinary specific, instead of re-protocling human ELISA kits and systems (Lane et al., 2018).

**Antibody Test Assays**

Antibody test systems are normally divided into two categories: those that test for the presence of an antibody in a biological sample, and those that use antibodies to test for chemicals in biological samples. One of the simplest tests that uses antibodies to detect chemicals present in biological samples is the lateral flow test kit, commonly used as a pregnancy test device. This kit can be used for early detection of conception onsite or in the field (Ambrose et al., 2007). However, with the massive increase in recent years in the range of ELISAs available for detecting and the quantification of molecules, most key biomolecules can be quantified using ELISAs. These systems vary enormously in sensitivity and the types of samples for which they are suitable, and the cost ranges from around £200 (US$283) for the commonly used/available kits to £1000 (US$1,417) or more for specialist kits.

ELISA assays are most commonly used in 96 well microplate format and normally allow around 80 samples (about 40 samples if performed in duplicate or about 24 samples in triplicate; see Figure 2) to be analysed. Up to 16 wells are required for calibration standards and controls. (These are important in order to confirm that the assay works correctly. They should not be skipped or reduced, however tempting it may seem.) As well as standard protocols, ELISAs can be modified for use in novel techniques—for example, in assessing chronic stress in dogs using cortisol as a biomarker extracted from nails as the biological source (Mack & Fokidis, 2017).

Sadly, the choice of ELISA is not always simple, as there are many different types. Generally, they can be classified into four main groups:

- **Direct ELISA:** Similar to the direct ELISA, but a second, conjugated antibody is used to detect the first unlabelled-antibody that bound the antigen.
- **Sandwich ELISA:** Antigen (target) is captured between two antibodies; one is bound to the plate and captures the antigen, and the other is added later for detection, thus forming a complex like a “sandwich.”
- **Competitive ELISA:** Measures the amount of sample by quantification of its interference/competition with an expected signal.

Potentially, the competitive ELISA is most useful for quantification of biomarkers because it allows accurate quantification of the levels present in biological samples (Cell Signaling Technology, n.d.), though it depends on the biomarker. Plotting calibration curves and calculating values for unknown samples can be time-consuming and potentially challenging. However, many free-to-use ELISA analysis websites will plot the
data and perform the calculations directly from the raw plate reader data (elisaanalysis.com).

**Enzyme Activity**

Unlike chemical and antibody assays, enzyme activity assays measure the activity of an enzyme, not just its concentration in a biological sample. Different isoforms of enzymes may vary in their rate of activity (Sulakhe & Lautt, 1987), so it can be important to measure how much usefully active enzyme is present. Enzyme activity is the key biological process causing modifications to other proteins, such as cleaving or changing sugars, in essence controlling and regulating all key responses within an organism. The activity levels of certain enzymes are altered with changes in an animal’s environment (e.g., quality of care or stress). In its simplest form, enzyme assays measure substrate cleavage; for example, amylase activity can be measured using the substrate blue starch. Amylase cleaves the sugar starch into smaller sugar molecules, releasing a blue dye that can then easily be measured. The amount of blue dye released is proportional to the amount of enzyme activity present within the sample (Abe et al., 1996).

However, as enzymes are an active biological molecule, their activity will change with temperature and the presence of certain chemicals. EDTA (ethylenediaminetetraacetic acid), a powerful chelating agent, sequesters ions such as calcium, iron, and zinc that are required by some enzymes for activity (Sabeur et al., 2001). So it is extremely important when designing enzyme assays to check that any buffer used for dilution is compatible with that enzyme assay.

**Polymerase Chain Reaction (PCR) Tests**

PCR tests amplify DNA from either a DNA or RNA template to aid in the identification of genes, mutations, or viral infection. These tests have a wide variety of uses, from identifying dogs that have mutations in the oxytocin receptor (OXTR) that can influence animal behaviour (Bence et al., 2017) to identifying species or breed types (Wu et al., 2018), and they can identify animals infected with a virus such as coronavirus (Chan et al., 2020). These PCR reactions take approximately 30 min to 3 hr to run. The reagent cost per reaction is comparatively inexpensive (starting around £0.30 [US $0.43]), but running the test requires a thermal cycler, which is a comparatively expensive piece of equipment.

RNA samples must be converted to DNA prior to PCR amplification, so an additional reverse transcriptase (RT) step is needed. This is important for identifying RNA-based viruses, such as those currently being used to detect COVID-19 infections in both humans and animals (Chan et al., 2020). The good news is that all-in-one RT-PCR kits are commercially available and can work from RNA in a single combined reaction step, but at a higher cost per sample.

**Proteomics**

Proteomics studies the variation in protein levels within a biological sample or even changes in the post-translational modification (PTM) status of proteins. Many factors increase/decrease protein levels to control processes in an organism, but at a cellular level, proteins can also be regulated once synthesised by PTM. For example, amino acid residues can undergo phosphorylation or acetylation within protein molecules. These PTMs can then alter the protein’s behaviour—for example, changing its affinity for a target receptor, changing the proteins it binds, and even changing its half-life (speed of degradation). Most proteomic analysis is through western blotting analysis using specialised antibodies that are specific to certain proteins, isoform specific antibodies, and even antibodies that only bind and recognise proteins with certain PTMs on specific amino acid residues/sites. Proteomics has even been used to identify changes in PTM/phosphorylation at a global protein level in saliva samples of dogs infected from suspected tick-borne babesiosis (Galán et al., 2018).

**Mass Spectrometry**

Mass spectrometry identifies the presence and relative amounts of chemicals and proteins in a biological sample. Mass spectrometry accurately determines the exact molecular mass of the molecules present in biological samples, and these are then compared with known standards. An example of its use is an investigative study on dogs treated with cannabidiol for canine epilepsy that used a simple and fast gas-chromatography mass spectrometry assay to accurately quantify cannabidiol metabolite levels in serum samples (Rotolo et al., 2019). Mass spectrometry is usually provided as a service, either academically or commercially, because of both the high level of expertise required to undertake the analysis and the cost of the equipment. Mass spectrometry (LC-MS/MS) can be extremely useful in identifying PTMs on proteins and small molecules in biological samples that cannot easily be determined by antibody methodologies (Koivunen et al., 2006).
Factors That Might Influence Selection of Biomarkers

Biological samples have a wide variety of influencing factors that are outside the researcher’s control. Therefore, these factors might influence biomarker or sample selection (Figure 1) and can be summarised as follows:

- Length of time the marker might be present
  - Half-life, excretion time, potential collection times (Wolff et al., 1999)

- Compounds that might affect readings
  - Medications or contamination, for example lysed blood samples will have falsely high levels of lactate dehydrogenase (Solberg et al., 1986)

- Medical conditions
  - For example, pancreatic disease can raise blood amylase levels, giving falsely high reading if using amylase is used as a marker of stress (Murtaugh & Jacobs, 1985)

The use of biomarkers to evidence animal cognitive, welfare, and health issues has vastly influenced the ability of a researcher to accurately and easily quantify biomarkers and directly correlate these to observed behaviours. In conclusion, no one-size-fits-all solution exists for the analysis of biomarkers; rather, the selection of sample type and biomarker is dependent on the biological samples available and the ability to analyse the biomarkers. Care should be taken when deciding on the sample to ensure that the results will answer the research question. The correct assay needs to be chosen to give the precise, required information. Making the wrong choices here can lead to confusing results and incorrect conclusions, and we all know how bad science can be propagated.

After all, the old saying rings true: Garbage in equals garbage out.

Biochemical Tips

- Extraction of saliva from swaps: Saliva can be extracted and collected in Eppendorf microcentrifuge tubes by cutting the swab about 5 mm from the end of the swab and inverting it so that the stick is at the bottom of the tube. The tube can then be centrifuged briefly (about 4-6000 xg for 10–30 s) to pellet the saliva at the bottom of the tube for easy pipetting.

- Dilution of samples: Some samples (e.g., saliva samples) might have to be diluted (in our experience, this is common for the larger breeds of dogs), as the viscosity of the saliva can be too high to allow for pipetting and analysis. Commonly these dilutions are around 1:4 with the assay buffer.

- More is better: Although it adds to the costs, it is always better to carry out the analysis more than once, so run a single sample on an ELISA plate in duplicate or triplicate. That way, any single anomaly is removed, and the results are more reliable. No assay is perfect, so there will always be some built-in errors, and this helps to reduce them (but never removes them).

- Keep your samples appropriately: If the samples need to be at 4 °C, −20 °C, or −80 °C, keep them there, and avoid freeze thawing on multiple occasions where possible, as this can affect the results. Although this takes up space, the samples are there to go back to if needed and can even be used for other studies, such as testing for antibodies in a pandemic. But make sure that your ethical approval states this.

- Remember safety comes first: For example, collection of saliva swabs from rabid dogs is never a good idea, but from ordinary dogs it is fine. Faecal samples often hide hidden pathogens and dangers such as Salmonella, which is zoonotic and can make people very ill.

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