Faecal biomarker and archaeobotanical analyses of sediments from a public latrine shed new light on ruralisation in Sagalassos, Turkey

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\textbf{Abstract}

A public latrine in the ‘Imperial Baths’ of Sagalassos was transformed into a dump site in the early Byzantine period. Several layers of urban waste, including ceramics, bones, glass and perhaps excrements were deposited on the floor. Faecal biomarker analyses and archaeobotanical analyses were conducted to reconstruct the history of the room. 5β-stanols of human origin, such as coprostanol, were found in the sewage channels together with mineralised plant remains, indicating a human faecal context. The botanical remains are furthermore representative of the Roman diet of the Sagalassians. Soil layers, deposited on top of the latrine floor and dating to the early Byzantine period, contained herbivore derived 5β-stanols, such as 5β-stigmastanol and epi-5β-stigmastanol. Additionally, a clear predominance of epi-5β-stanols over 5β-stanols showed that the animal dung has been subject to composting. In this period, the former latrine was clearly used as a manure production site which is further confirmed by stratigraphic evidence of large amounts of urban waste artefacts, which were commonly collected together with manure before application on the fields. The results of the present study support the theory that off-site potsherds scattering can be used as a proxy for manuring events. Additionally, the data show key evidence for vertical migration of 5β-stanols and presumably also for the leaching of bile acids.

\section{Introduction}

Latrines, cesspits and other types of waste repositories provide a wealth of information on cultural, economic and environmental aspects of ancient civilisations. Apart from human or animal excrements, such contexts retain many artefacts reminiscent of ancient daily life such as pottery, food refuse, fodder, glass and coins (e.g. De Groote et al., 2004; Deforce, 2010; Greig, 1981; Knights et al., 1983; Tyson, 1996). Faecal remains and waste deposits can be studied by several approaches, dealing either with the identification of faecal matter in relation to the archaeological context, or to the taphonomic and diagenetic processes that affected the in situ preservation. Initially, mainly palaeobotanical analyses, i.e. the analysis of macrobotanical remains such as plant leaves, seeds and stems as well as pollen analysis, have been performed on such deposits (Greig, 1981, 1982). There is also a large potential for their systematic microscopic investigation in order to elucidate questions on ancient trade, farming or gardening, diet, cooking and the environment (Deforce, 2010; Martinez, 2005; Oeggl et al., 2009; Ramsay and Tepper, 2010).

Another approach is the microscopic study of soils in thin sections, or micromorphology, which can provide identification of dung layers through observation of opal phytoliths, dung spherulites, calcium minerals and other dung related features (MacPhail et al., 2004; Matthews, 2010; Shahack-Gross and Finkelstein, 2008; Shillito et al., 2011b). Study of the taphonomic pathways affecting the preservation state of these features can provide information on herding strategies, transhumance, subsistence and other socio-economic themes (Shahack-Gross, 2011). Inferences on the health status of ancient populations can be made by studying intestinal parasites (Reinhard, 1992; Mitchell and Tepper, 2007). The origin and diagenetic fate of dung-derived C and N pools is
investigated by stable isotope analysis (Dungait et al., 2009, 2010; Ghosh et al., 2003; Shahack-Gross and Finkelstein, 2008; Simpson et al., 1999a). Analysis of phosphate, one of the main indicators of human activity, and multi-element analyses are interpretative tools to elucidate patterns of human occupation and cultivation, such as the identification of middens, burial soils, cultivated farmlands and byres (Degryse et al., 2003; Entwistle et al., 2000; Farswan and Nautiyal, 1997; Parnell et al., 2001; Simpson, 1997). A more rigorous identification of faecal matter can be achieved by analysing faecal biomarkers (e.g. Bethell et al., 1994; Knights et al., 1983; Pepe et al., 1989; Pepe and Dizabo, 1990). These markers are organic compounds derived from a faecal source, which retain their chemical structure during diagenesis (Bull et al., 2002). In addition to the identification of faecal material, they also provide information on their biogenic origin.

A first class of faecal biomarkers are 5β-stanols. Their presence in ancient coprolites and waste repositories has evidenced their persistence over several millennia, which enables their use in archaeological studies (Bull et al., 2005; Knights et al., 1983; Lin et al., 1978; Pepe et al., 1989; Pepe and Dizabo, 1990). Their presence in coprolites of extinct mammals even demonstrates their use in Pleistocene research (Gill et al., 2009; van Geel et al., 2008, 2011). In addition, it has been experimentally confirmed that 5β-stanols persist in soils which received manure during a prolonged period (Bull et al., 1998; Evershed et al., 1997). Another group of faecal biomarkers are bile acids. Like 5β-stanols, bile acids have been shown to persist over several millennia in archaeological faecal deposits (Lin et al., 1978; Bull et al., 2005). Some authors even argue that they are more recalcitrant than 5β-stanols (Bull et al., 2003; Ehmmali et al., 1997, 2000). Both groups of faecal biomarkers have been widely applied in archaeological studies. Human faeces, for example, have been identified in ancient cesspits, middens and latrines (Bethell et al., 1994; Bull et al., 2005; Pepe et al., 1989; Pepe and Dizabo, 1990; Shillito et al., 2011a, 2011b), in a Roman fortification ditch (Knights et al., 1983) and in sediments from a Roman street drain (Bull et al., 2003). Also, the use of herbivore or porcine dung as manure for agricultural soils has been investigated (Birk et al., 2011; Bull et al., 1999, 2001; Evershed et al., 1997; Simpson et al., 1999b). Recently, archaeol (a dialkyl glycerol ether) has been proposed as a biomarker for forage fermentation in ruminants (Gill et al., 2010).

Numerous latrines from the Roman and Medieval period have been studied in Western Europe. Evidence from the Eastern Mediterranean and especially Anatolia, however, is still scarce. The current paper focuses on the analysis of latrine deposits from the Roman to early Byzantine town of Sagalassos, south-western Turkey. The stratigraphy of the sediments provides indications for three use-phases. Architectural remains evidence that the room functioned as public latrine during its first phase. The functionality during later phases, however, is still unclear. Its use during Early Byzantine Sagalassos (mid 5th – 7th century AD) is of particular interest as this period was a turning point in the city's history, which was marked by increasing ruralisation and ultimately urban decline. A multi-analytical approach, using analysis of faecal biomarkers, calcium analysis, macrobotanical remains, pollen and fungal spores, was chosen in order to achieve a more precise understanding on the use and maintenance of the studied structure. Particularly the combination of techniques has proven to overcome the limitations of each individual approach (e.g. Shillito et al., 2011b). More specifically, this paper aims at (i) identifying the presence and source of faecal remains, (ii) analysing the general organic (and inorganic) content of the structure and (iii) integrating these data to elucidate the different use-phases and associated maintenance policies of the structure.

2. Study area

2.1. The site and its importance

Sagalassos is located in southwest Turkey, in the Pisdian Lake District, at an altitude of 1450–1600 m a.s.l. in the Western Taurus mountains, approximately 110 km north of Antalya (ancient Attaleia) (Fig. 1). The natural assets of the territory, its strategic location, abundant water, fertile soils, availability of a variety of raw materials, and a major road connecting the city with the harbours on the Anatolian south coast fuelled the development of the city as a regional centre (Vanhaverbeke and Waekens, 2003). Under Roman Imperial rule (from ca. 25 BC – mid 5th century AD) the city flourished and became the most important town in Pisidia. The late Roman period (300–450 AD) witnessed the most intense exploitation of the city's territory. The first 100 years of the early Byzantine period (mid 5th – 7th century AD) were still a period of prosperity in both city and countryside. Yet, over the course of the 5th century, signs of stress on the city's subsistence patterns appear together with more intensive farming in the immediate vicinity of the city (Degryse et al., 2004). From the mid 6th century AD onwards, there is increasing evidence of a ruralisation of Sagalassos, possibly coinciding with a decrease of the urban population and the probable abandonment of some quarters. From the onset of the excavations in 1990, attention was paid to monumental and architectural features as well as to interdisciplinary research, in order to better understand archaeological contexts in view of functional and social meaning, raw materials and the way they are treated, subsistence, human impact on the landscape and the palaeoenvironment.

2.2. Description of the latrine

The studied profiles originate from one of the interconnected vaulted rooms supporting the imperial bath complex, built ca. 120–165 AD, i.e. room 4 (Fig. 2). This room measures approximately 8 by 10.5 m and has two windows. At the time of its discovery, room 4 was filled almost to its ceiling with mortar and small brick fragments from the covering of its concrete ceilings. The stratigraphy of the room was very complex and was described in more detail in previous records (Vermoere et al., 2003, 2004). The presence of channelled stones at the original floor level and holes for fixing seats made it clear that the room had originally served as a public latrine (Fig. 2). The sewage channels, constructed alongside the room, served to collect human excrement and to drain them into the entrance of a sewer in the north-western corner of the room. The room was supplied with water which was recycled from the baths at the upper floor level and supplied to the room through a channel in the eastern wall (Fig. 2). The structure thus represents one of the best surviving examples of Roman greywater reuse.

At a certain point in the history of the room, as discussed below, the toilet seats were removed and the room served as a collector of waste material. Sediment layers excavated above the original floor level – part of the stratigraphy is still present at locus A (Fig. 2) – contained a large amount of urban debris such as glass, coins, marble slabs, tessereae, metal, ceramics and animal bones, dating to the mid-3rd–6th centuries AD. The fine ware ceramics found near the original floor level displayed traces of water abrasion and surface trampling, suggesting that the material was collected elsewhere and intentionally dumped here. Most of the bone material represented consumption refuse including bones from ovicaprines, cattle, pig, chicken and hare. Pollen analysis of these sediment layers complying with the stratigraphic evidence that waste material was collected from different parts of the city (Vermoere et al., 2003, 2004). Mineralogical analysis by X-ray
diffraction (XRD), conducted on the same sediment layers, furthermore indicated the possible addition of lime as revealed by the presence of mainly fine-grained calcite (CaCO$_3$) and minor amounts of portlandite (Ca(OH)$_2$) and lime (CaO) (Patrick Degryse, pers. comm. 2008). In addition, the simultaneous occurrence of brushite and weddelite in certain layers could suggest a urinary origin (Daudon et al., 1998; Estepa and Daudon, 1997), although these minerals might also originate from other sources or have been formed by mineralisation of organic material (McCobb et al., 2001, 2004). In the western part of the room, the floor level was raised twice in the course of the 7th century AD (within a short period), demonstrating that this part of the room was once more in regular use. At the time of discovery, the room was completely filled with mortar and small brick fragments.

3. Material and methods

3.1. Coring and sampling of sediments

Core profiles were taken at different locations within room 4, with sub-samples taken every 10 cm. Three cores of about 60 cm depth each were taken from the stratified sediment heap that remained from the 1998 excavations (locus A, Fig. 2). Another two cores of 140 cm and 180 cm depth were taken using an Edelman corer from the sewage channel at loci C and E, respectively. The sediments were divided into categories as depicted in Fig. 2. Soil pH was determined by suspending 5 g of soil into 25 ml of 0.01 M CaCl$_2$ and measuring the pH of the resultant slurry. Six to seven samples were analyzed per locus (Table 1). A summary of all sampled sediments is listed in Table 1 along with the stratigraphic description and applied methods. Charcoal fragments were isolated from four sub-samples for obtaining radiocarbon dates. The $^{14}$C ages were processed with accelerator mass spectrometry (AMS) after a standard acid-alkali-acid pre-treatment. The radiocarbon dates (Table 2) are quoted in conventional $^{14}$C years by correction for isotopic fractionation using $\delta^{13}$C values, and are calendar calibrated dates according to the INTCAL04 database (Reimer et al., 2004).

3.2. Faecal biomarker analyses

Extraction and purification of faecal biomarkers is based on a method described by Bull et al. (1999, 2001). We modified this procedure by using liquid–liquid extraction of the saponified lipid extract instead of the solid phase extraction to separate stanols and bile acids. This was achieved by extracting neutral lipids (including stanols) with hexane (2–3 times). The water content was then raised to 60% and after acidification (6 M HCl, pH 3–4), polar lipids...
(including bile acids) were extracted with ethyl acetate (2–3 times). The solvent of both fractions was removed under a gentle stream of nitrogen. Subsequently, both fractions were derivatised with N-methyl-N-(trimethylsilyl)trifluoroacetamide (MSTFA) with 1% trimethylchlorosilane (TMCS) at 60 °C for 1 h yielding trimethylsilyl derivatives. After removal of solvent and excess reagent, neutral and polar fractions were redissolved in hexane and ethyl acetate, respectively. Both fractions were analysed with gas chromatography coupled to mass spectrometry (GC–MS) using an Agilent 6890 N GC instrument, equipped with a HP5-MS capillary column (30 m × 0.25 mm i.d. × 0.25 μm film thickness) and coupled to an Agilent 5973 Network Mass Selective Detector. One μl of each fraction was injected in the splitless mode at a temperature of 290 °C. The oven temperature was held at 150 °C for 2 min, increased to 250 °C at 20 °C/min followed by an isothermal 10 min hold and further ramped to 300 °C at 2 °C/min followed by an isothermal hold of 20 min and a final ramp to 340 °C at 10 °C/min. The mass spectrometer was held at a temperature of 340 °C and operated in scan mode (spectra were recorded between m/z 50 and 700) and in selected ion monitoring (SIM) mode. Peak

**Fig. 2.** Map of the ground floor of the bath complex of Sagalassos with location and stratigraphic description of the core profiles in room 4 and 5.
assignments were made by comparison with retention time of authentic standard compounds and literature mass spectra. Reliable identification and quantification was achieved by adopting GC–MS/SIM as analytical technique. The diagnostic mass fragments used to detect target sterols and stanols are listed in Table 2.

3.3. Measurement of Ca concentration by elemental analysis

The soil samples were first dried and finely ground. Calcium was then extracted in duplicate by digestion of 0.2 g soil material in 5 ml of a mixture of 70% HClO₄, 65% HNO₃ and 37% HCl (1:3:1 v/v/v) in a microwave system (Milestone MLS 1200). Samples were subsequently diluted and concentrations of calcium were determined by atomic absorption spectroscopy (AAS, Solaar 969 AA).

3.4. Macrobotanical analysis

The average volume of the samples taken for the analysis of plant macrofossils comprised ca. 50 cm³. The samples were mixed with water for 12–24 h in order to gently suspend them. The plant macrofossils were then recovered by wet sieving using sieve meshes of 1 mm and 0.3 mm. The plant materials were sorted using a binocular microscope (Zeiss Stemi 2000 CS) and identified through comparison with the reference collections of the Centre for Archaeological Sciences, Katholieke Universiteit Leuven. The absolute numbers of counted plant remnants per sample depths are represented in Table S2 and plotted in relation to the results of the biomarker analyses in Fig. 4.

Table 1
Summary of sampled sediment layers per locus with their stratigraphic category and applied methods. The location of the sediment cores is indicated in Fig. 2.

<table>
<thead>
<tr>
<th>Sample number</th>
<th>Depth (m)</th>
<th>Stratigraphy</th>
<th>pH</th>
<th>Faecal biomarkers</th>
<th>Ca</th>
<th>Macroremains</th>
<th>Pollen</th>
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<td>SA08CV1263</td>
<td>[0.0; 0.1]</td>
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<td></td>
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<td></td>
<td></td>
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<td>7.5</td>
<td></td>
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<td>2</td>
<td>7.5</td>
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<td>2</td>
<td>7.5</td>
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<tr>
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<tr>
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<td>2</td>
<td>7.5</td>
<td></td>
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<td></td>
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<tr>
<td>Locus C – sewage channel in northeastern corner</td>
<td></td>
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<td>7.5</td>
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<td>7.5</td>
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<td>Locus E – sewage channel in middle of northern wall</td>
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<td>7.5</td>
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<td>Locus A – heap close to southwestern window</td>
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Table 2
AMS 14C dates of the latrine deposits.

<table>
<thead>
<tr>
<th>Sample number</th>
<th>Locus</th>
<th>Depth (m)</th>
<th>Laboratory code</th>
<th>δ¹³C (‰ PDB)</th>
<th>Conventional ¹⁴C age (BP)</th>
<th>Cal. (2σ) age</th>
<th>Cal. (1σ) age</th>
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<td>A</td>
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<td>Beta-281735</td>
<td>-24.1</td>
<td>1550 ± 40</td>
<td>AD 420–600</td>
<td>AD 430–560</td>
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<tr>
<td>SA08CV1265</td>
<td>C</td>
<td>[0.2; 0.3]</td>
<td>Beta-284661</td>
<td>-22.0</td>
<td>1550 ± 40</td>
<td>AD 420–600</td>
<td>AD 420–560</td>
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<td>SA08CV1276</td>
<td>E</td>
<td>[1.1; 1.4]</td>
<td>Beta-281733</td>
<td>-24.9</td>
<td>1730 ± 40</td>
<td>AD 230–410</td>
<td>AD 250–380</td>
</tr>
<tr>
<td>SA08CV1293</td>
<td>E</td>
<td>[1.6; 1.7]</td>
<td>Beta-281734</td>
<td>-22.2</td>
<td>1780 ± 40</td>
<td>AD 130–350</td>
<td>AD 220–320</td>
</tr>
</tbody>
</table>

Notes:
- Depth is expressed relative to the original floor level of room 4 (negative depths reflect sediments beneath floor level).
- Numbers refer to categories as explained in the text and in Fig. 2.
- Charcoals fragments from these sediment layers were taken for ¹³C dating (see Table 2).

3.5. Analysis of pollen and fungal spores

Samples of 3 cm$^3$ were taken every 10–15 cm for pollen analysis. The samples were processed according to Faegri and Iversen (1989). Clay contamination was removed by ultrasonic sieving (5 μm) although some pollen grains (Juniperus) could be lost by this procedure. Nevertheless, up to 250 pollen grains of terrestrial plants were counted. The pollen sum consists of terrestrial arboreal (AP, trees and shrubs) and non arboreal pollen (NAP, herbs) and excludes aquatic and marsh plants. Non-Pollen-Palynomorphs (NPP) were counted in the samples prepared for routine pollen analysis. The “types” identified, follow the type numbers described by van Geel (1978, 1986, 2001) and van Geel et al. (1995, 2003). For calculation and diagram construction the programmes TILIA and TILIA GRAPH were used (Grimm, 1991). The pollen diagram (in percentages) is depicted in Fig. 6.

4. Results and discussion

4.1. Sediment description and radiocarbon dating

In terms of stratigraphy, locus A is situated above loci C and E and comprises 14 layers, subdivided into 6 different categories (Fig. 2, Table 1). Category 1 comprises a heterogeneous reddish brown silt layer with charcoal, volcanic tuff and large ceramic fragments, and can be related to a dump context. The yellowish clay layers of category 2 serve as a key horizon and together with the sediments of category 3 they can probably be associated with lime addition (see also Section 4.3). Categories 4 and 5 represent moister phases of activity (see also Section 2.2) and the black organic rich clay layers (category 6) represent dry periods following inundation.
Fig. 4. Combined results of faecal biomarker analysis, elemental (Ca) analysis and analysis of macrobotanical remains of the latrine sediment cores. Depths are expressed relative to original floor level with positive and negative values reflecting samples above and beneath floor level, respectively. Sediment dates are expressed as the 2 sigma calibration interval resulting from radiocarbon dating. The location of the different soil profiles is indicated in Fig. 2. Labelled compounds represent coprostanol (1a), epicoprostanol (2a), cholesterol (1c), 5α-cholestanol (1d), 5β-stigmastanol (3a), epi-5β-stigmastanol (3b), sitosterol (3c) and 5α-stigmastanol (3d). Chemical structures of the steroids are given in Fig. S1.

Fig. 4. (continued).

Fig. 5. Summary of the stanol ratios for the latrine sediments: (a) \([5\beta + 5\alpha] C27 + 29\) stanol \([\square = C_{27}; \bigcirc = C_{27-29}\) stanol] ratio of \([\square = C_{27}; \bigcirc = C_{27-29}\) stanol], (b) \([\text{coprostanol}: 5\beta - \text{stigmastanol}] \) ratio (\([\vartriangle = 3\alpha\)-epimers excluded \(\bigcirc = 3\alpha\)-epimers included] and \((\text{coprostanol} + \text{epicoprostanol}): (5\beta - \text{stigmastanol} + \text{epi-}5\beta - \text{stigmastanol})\) ratio (\([\vartriangle = C_{27}\) stanol \(\bigtriangledown = C_{27}\) stanol]), (c) \([\text{coprostanol}: \text{epicoprostanol}] \) ratio (\([\bigcirc = C_{27}\) stanol] and \([\text{epi-}5\beta - \text{stigmastanol}: \text{epicoprostanol}] \) ratio. Depths are expressed relative to original floor level with positive and negative values reflecting samples above and beneath floor level, respectively. The location of the soil profiles is indicated in Fig. 2. Chemical structures of the steroids are given in Fig. S1.
At loci C and E, ca. 30 cm of debris accumulation was found on top of a 90 cm thick light grey clay layer (Fig. 2). Below, at least 60 cm of dark grey organic rich clay is present (Fig. 2). Both clay layers contain a large number of small charcoal, ceramic and volcanic tuff fragments. From 30 cm onwards, the profiles of loci C and E are hypothesized to be related to latrine deposits, while the upper sections of profiles C and E and the profile at locus A may be linked to the post-latrine use of the room.

The pH values of the sampled sediment layers are listed in Table 1. The pH range was rather narrow with values between 7.5 and 8.0. No significant differences between the different loci were observed.

Four samples of the different latrine deposits were chosen for AMS radiocarbon dating (Table 2). The dates displayed an orderly relationship with depth, irrespective of the location of the samples. Two samples, taken from near the bottom of the sewage channel at locus C (depth of 130 cm within the dark grey clay layer, Fig. 2) and E (depth of 160 cm within the dark grey clay layer, Fig. 2), displayed an earlier date of AD 230–410 and AD 130–350 respectively, corresponding with the first phase of room 4, namely a public latrine. The latter date demonstrates that the latrine was already in use shortly after the construction of the ‘Imperial Baths’ (120–165 AD) and was not a later 6th century AD transformation as the excavators originally assumed (Waelkens et al., 2006). The remaining two samples, one from the stratified sediment heap that remained from the 1998 excavation (locus A, between 0 and 15 cm above original floor level) and another from the top level of the sewage channel (at 20 cm depth within the anthropogenic waste layer at locus C), showed equal calibrated (2σ) dates of AD 420–600, corresponding with the second phase of room 4. These observations match the stratigraphic observations in Section 3.1.

### 4.2. Faecal biomarkers

The total lipid extracts of the latrine deposits contain a diverse set of lipids, including fatty acids, alkanols, alkanes, steroidal lipids, diterpenoids and pentacyclic triterpenoids. For the purpose of this study, only steroidal lipids, viz. sterols, stanols and bile acids, will be discussed.

**4.2.1. Evidence for faecal derived 5β-stanols**

The presence of 5β-stanols, such as coprostanol and 5β-stigmastanol, in the latrine deposits provides a first indication of faecal matter, as they are products of the enteric hydrogenation of Δ4-sterols (Björkhem and Gustafsson, 1971). Epi-5β-stanols, such as...
epicoprostanol and epi-5β-stigmastanol, were also present, but are normally not abundantly present in fresh faeces (Leeming et al., 1996). They can however be found in soils as a result of microbially mediated epimerisation of 5β-stanols (Mackenzie et al., 1982; McCalley et al., 1981). Unlike 5β-stanols, 5α-stanols are not faecal biomarkers but are primarily formed by Δ5-sterol reduction under aerobic conditions (Gaskell and Eglinton, 1975; Taylor et al., 1981).

As 5β-stanols in sediments may also derive from a non-faecal origin, for instance as a result of sedimentary reduction of sterols, the interpretation of absolute concentrations of individual stanols is questionable (Bull et al., 2002). Therefore, the [5β:(5α + 5x)] stanol ratio has been established by studying modern-day sewage pollution (Grimalt et al., 1990). The value of 0.7 has been proposed as a lower limit for faecal pollution whereas values between 0.1 and 0.3 are representative for non-contaminated sediments. This ratio was further adapted by Bull and others (1999) to account for microbial and diagenetic changes that may affect the relative concentration of these biomarkers through time (Mackenzie et al., 1982; McCalley et al., 1981):

\[
\text{[(5β-stanol + epi-5β-stanol) : (5β-stanol + epi-5β-stanol + 5α-stanol)]}
\]

(1) The values of this stanol ratio for the latrine sediments were plotted in function of depth, as depicted in Fig. 5a. The values have been calculated for both C27 stanols and C27β:29α stanols, as ruminant dung is relatively enriched in C29 stanols such as 5β-stigmastanol (cf. infra) (Bull et al., 1999; Leeming et al., 1996). Values of the C27 stanol ratio (1) above 0.7 were clearly observed in the sediments of locus E and unambiguously demonstrate the faecal nature of the 5β-stanols. The C27α:29β stanol ratios were notably lower in the top layers of this profile due to the enhanced concentration of 5α-stigmastanol (Fig. 4) while a rapid increase to 0.7 was observed at depths below 60 cm. At locus C, deep layers (30–140 cm) also displayed elevated values, ranging between 0.51 and 0.76. However, both the upper layers (0–30 cm) from this core profile and the sediments from locus A displayed distinctly lower C27 and C27α:29β stanol ratios, with values around 0.45 and 0.55 respectively. These intermediate stanol ratios do not meet the faecal threshold of 0.7 as proposed by Grimalt et al. (1990). However, as Birk et al. (2011) have recently pointed out, this threshold was determined for urban sewage pollution in coastal waters and cannot be applied in general for archaeological sediments. Firstly, sedimentary reduction of Δ5-sterols can give rise to enhanced concentrations of 5α-stanols due to the higher thermodynamic stability of the latter compared to 5β-stanols (Mackenzie et al., 1982; van Graas et al., 1982). Secondly, deposition of organic waste or litter fall would effect a rise in 5α-stanol concentration and, consequently, a dilution of faecally derived 5β-stanols. Hence, the [5β:(5α + 5x)] stanol ratio in ancient sediments can be biased by diagenetic and post-depositional factors. Indeed, studies on Roman cesspits and ancient anthropogenic sediments report values of the stanol ratio (1) below 0.7 (Bethell et al., 1994; Bull et al., 1999, 2001).

Nevertheless, these arguments do not explain the discrepancy between the high values (above 0.7) of locus E on the one hand, and the intermediate values (around 0.5) of locus A and the top layers of locus C on the other hand. The dissimilar nature of these sediments was already observed in their stanol profiles: while animal-derived C27 stanols were the major steroids in the former, plant-derived C29 stanols clearly predominated in the latter (Fig. 4). As herbivores ingest large amounts of plant matter, the predominance of C29 5β-stanols can be attributed to the presence of herbivore dung (Bull et al., 1999, 2001). This is further endorsed by the fact that herbivore dung is typically enriched in 5α-stigmastanol concentrations leading to intermediate stanol ratios (Evershed et al., 1997; Leeming et al., 1996). By contrast, omnivore excreta are characterised by a dominance of C27 5β-stanols and values of stanol ratio (1) above 0.7. This was clearly observed in the sediments of locus E as well as in some of the deep layers of locus C. In the latter, a slightly decreased C29 stanol ratio (1), with values between 0.6 and 0.7, was observed between 50 and 120 cm depth. In the following section, the biogenic origin of the 5β-stanols will be discussed in more detail.

4.2.2. Evidence for human and animal 5β-stanols

As herbivores ingest large amounts of plant-derived sterols, their excreta are enriched in C29 and C29 5β-stanols such as 5β-campestanol and 5β-stigmastanol (Leeming et al., 1996). Therefore, the profile of faecal 5β-stanols allows for discrimination between herbivores and omnivores. In the light of this observation, the C27/C29 5β-stanol ratio was proposed by Bull et al. (2001) and Evershed and Bethell (1996):

\[
\text{coprostanol : 5β-stigmastanol (2)}
\]

This ratio can be used as a proxy for the biogenic origin of 5β-stanols with values for humans and pigs ranging between 1.5 and 5.5, and values for ruminants being in the order of 0.25.

In Fig. 5b values of this stanol ratio are plotted in function of sample depth. Ratios of the core profile from locus E ranged from 1.06 to 3.45 with only shallow samples (0–30 cm) exhibiting values below 1.5. The high values in the deep soil layers (30–190 cm) clearly indicate an omnivore origin of the 5β-stanols. By contrast, both locus A and the top levels (0–30 cm) of locus C exhibited very low C27/C29 stanol ratios around 0.05, thus confirming a herbivore origin of the stanols in these sediments. The deep layers (30–140 cm) of locus C displayed values of the stanol ratio (2) between 1.0 and 1.5.

4.2.3. Evidence for composting of faecal matter

Another striking observation is the higher abundance of epi-5β-stigmastanol compared to 5β-stigmastanol at locus A and in the shallow soil layers (0–30 cm) of locus C (Fig. 4). The same pattern was noted for coprostanol and epicoprostanol at locus A. This can be illustrated by plotting the 3β/3α-stanol ratio in function of depth (Fig. 5c):

\[
\text{coprostanol : coprostanol (3)}
\]

McCall et al. (1981) have shown that C3β,5β-stanols can be converted to their 3α-epimers during reworking in anoxic environments. Therefore, low 3β/3α-stanol ratios (<1), as observed for locus A and for the shallow samples (0–30 cm) of locus C, indicate that the excreta have been subjected to composting. Similar observations were noted in Minoan terraces from Crete (Bull et al., 2001). In this study, the enhanced abundance of 3α-epimers was interpreted as arising from predepositional reworking of faecal matter as the dry and arid conditions of soil formations of Crete are unlikely to promote such reactions.

In contrast, the omnivore excreta found in the sewage channel at locus E were not subjected to a similar composting process as the 3β/3α-stanol ratio (3) displayed only values above 1.0. A similar pattern was observed for the C27 stanol ratio (3) of locus C. In contrast, the latter core profile exhibits a gradual increase of the C29 stanol ratio (3) from values around 0.5 in the top layers towards values around 3.1 in the bottom layers. The latter will be further discussed in the following section.

4.2.4. Indications for vertical migration of stanols

The aforementioned stanol ratios unambiguously evidence the presence of composted ruminant dung at locus A and in the upper
layers of the sewage channel at locus C, and of non-composted omnivore faeces in the sewage channel at locus E. The deep layers (30–140 cm) of locus C, nevertheless, still exhibit some ambiguity. These layers exhibited a markedly distinct stanol profile in comparison with the superimposed herbivore dung layers (cf. Fig. 4), but also exhibited a number of dissimilarities with the stanol profile of locus E. The relative concentration of Δ5-sterols, for instance, was generally higher in locus C. Additionally, C27 and C29 stanols were almost equally abundant in locus C, whereas locus E displayed a marked predominance of C27 stanols. This is surprising since both core profiles were taken from the same sewage channel and were only about 4 m located from each other. Similar ambiguities were found in the stanol ratios of the deep layers (30–140 cm) of locus C. The [5β:5β+5α] stanol ratio (1) exhibited values between 0.6 and 0.7. These values are situated between the values for herbivore and omnivore dung, as observed in locus A and E, respectively. The C27/C29 stanol ratio (2) also furnished intermediate values close to the empirical threshold of omnivore faeces. A possible explanation would be that both herbivore and omnivore excrements were deposited in the sewage channel in the same period. However, the gradual increase of the C29 stanol ratio (3) suggests a second hypothesis, namely that stanols of the herbivore dung layers (0–30 cm) of locus C infiltrated down the sewage channel and intermingled with stanols of omnivore excrements. This can be further illustrated by plotting the ratio of epi-5β-stig 

\[ \text{5β-stigmastan-3z-ol : 5β-cholestan-3z-ol} \] 

(4)

This stanol ratio (4) exhibits a marked inverse proportionality with depth. The data points were fitted with a power regression and yielded a correlation coefficient of 0.89. Such a gradual concentration pattern is unlikely to result from a mixed deposition with depth. The data points were presumably located east of this profile. The latter is possible since water entered the room through a channel in the eastern wall and flowed into the sewage channel towards the sewer entrance in north-western corner of the room (Fig. 2).

4.3. Calcium analysis

Several yellowish layers became apparent from the stratigraphy of the sediment cores, particularly in the soil profile at locus A (see photograph and stratigraphic category 2 in Fig. 2). Because of the possibility of lime addition it was decided to measure elemental concentrations of calcium. The results are depicted in Fig. 4. At medium depths (30–140 cm) in the sewage channels (loci C and E), the calcium concentration exhibited low values, namely between 50 and 80 mg Ca/g soil. Conversely, the shallow soil layers from both cores and the soil layers from locus E below 1.5 m tended to be more enriched (up to 146 mg Ca/g soil). All samples from locus A exhibited an even higher calcium concentration around 190 mg/g soil (Fig. 4).

4.4. Macrobotanical analysis

The macrobotanical content of the sewage channel cores consisted predominantly of charred and mineralised plant remains. The co-occurrence of these types of plant remains indicates rather complex depositional conditions. They could have become incorporated into the deposits by flushing and subsequent perturbation of the sewage channel. The charred remains, including cereals (e.g. *Triticum* sp., *Hordeum vulgare*), *Panicum miliaceum*, lentils (*Lens culinaris*) and walnuts (*Juglans regia*), most probably derive from ashes, which were either thrown into the sewage channel as desiccant to prevent bad smell or discarded therein as waste. Of greater interest are the mineralised plant remains. Mineralisation can occur in archaeological contexts when organic matter is exchanged by calcium phosphate (McCobb et al., 2001, 2004). Roman latrines frequently exhibit such preservation conditions as calcium phosphate is an important constituent of faeces (Jacomet and Kreuz, 1999). In addition to charred and mineralised plant remains, also fragments of strongly decomposed organic matter was present, particularly in the lower part (70–170 cm) of locus E. Although these fragments could not be identified, the presence of such highly decomposed matter suggests that this part of the sewage channel was subject to water logging.

The identified mineralised seeds in the sewage channel profiles, *viz.* fig (*Ficus carica*), plum (*Prunus* sp.) and grape (*Vitis vinifera*), constitute a number of frequently recorded species in the Roman food chain (Kuijper and Turner, 1952; White, 1970) and were also quite common in Sagalassos during the Roman and early Byzantine period. Most of these remains are primarily used for human consumption rather than for fodder, and thus make up an important part of the diet of the Sagalassians from the 2nd to 5th century AD. Notably, pits of olives, which represent an important element of Roman agriculture and diet, were not found in the latrine deposits. This is remarkable since other fruit stones, such as *Prunus* sp., were present and would generally preserve in a similar manner as olive pits. Most probably, olives were not a very common component in the diet of the Sagalassians as archaeobotanical analyses of Roman to early Byzantine urban contexts show that charred olive stones only account for ca. 3% of the total amount of investigated plant remains. Finally, mineralised remains of spices were also found in the lower part for locus C (*Table S2*), such as the inner part of probably coriander and dill. Both condiments were rather popular for Roman food preparation (Prance and Nesbitt, 2005 and references therein). Their occurrence and preservation in mineralised state is strong evidence for the presence of human faeces in the studied sediments and correspond to the first phase of room 4, namely a public latrine.

Apart from plant macroremains also some small animal remains were found. However, only few of them could be identified and included one bone of a micro-mammal and six fish remains, i.e. 1 rib, 3 scales and 2 pharyngeal plates from very small cyprinids (130–140 cm deep at locus C and 160–170 cm deep at locus E). All other fragments (66 in total) are small pieces of bird and mammal bones, which remained unidentifiable. Traces of digestion (or the effect of gastric juices) are not clearly visible on these bones. It is therefore questionable but not excluded that the bones have passed through the digestive tract (Bea De Cupere, pers. comm. 2010).

4.5. Pollen analysis and fungal spores

Pollen preservation of the studied profiles was mainly limited to the lower part (70–170 cm) of locus E. The pollen found, moreover,
generally represent the vegetation in the surrounding of the sampled location and are to a much lesser extent representative of human food (Fig. 6). This is most probably due to an influx of local and regional pollen with water percolating through the sewage channel. Several of the pollen taxa, belonging to the group of “cultivated” species, originate from trees that most probably grew in the city, such as chestnut (Castanea), hazel (Corylus), walnut (Juglans) and plum (Prunus). Finds of hazel, walnut and plum are common vegetation elements of this site as they appear also in wood charcoal assemblages from the Early Byzantine period. These species were furthermore exploited for alimentary purposes (see Romanus et al., 2009).

4.6.1. The first phase: a public latrine

The first phase of room 4 is clearly associated with the function of public latrine, as pointed out by the architecture of the vaulted room (see Section 2.2). The construction enables the cleaning of the channel by flushing periodically with large amounts of water that was recycled from the baths on the upper floor levels. This is also visible in the pollen composition of the sewage channels, representing the wild growing vegetation, predominantly woodland and open vegetation, in the surrounding of the ‘Imperial Baths’ (Fig. 6). Nevertheless, water logging must also have occurred, at least partly, as suggested by the presence of strongly decomposed organic matter in the sewage channel profile at locus E. This event, however, may also have happened after the function of latrine was abandoned.

The radiocarbon dates of the lowest sample of the sewage channel at locus E (130–350 cal AD) showed that the latrine was already in use shortly after the construction of the baths in 120–165 AD. Conclusive evidence for faecal remains was provided by the presence of faecal biomarkers, viz. 5β-stanols, in the sewage channel (loci C and E). The stanol ratios of the deep layers are rather uniform and thus confirm that no significant change in faecal deposition has taken place. The [5β:(5β + 5α)] stanol ratios in the deep sediments of the sewage channel cores displayed values between 0.6 and 1.0 and were interpreted as deriving from faecal matter (Fig. 5a). Despite recent criticism on the applicability of this stanol ratio for the assessment of ancient manuring events (Birk et al., 2011), these values sustain the validity of this ratio in concentrated faecal deposits such as latrines. The coprostanol:5β-stigmastanol ratio of these layers exhibits values above 1.5 and indicates an omnivore faecal deposition, most probably from humans (Fig. 5b). Apart from faecal biomarkers, the sewage channel also contained traces of consumption refuse. These include macroscopic seeds of cereal grains, lentils, beans, figs, walnuts, plums, grapes, and inner parts of spices, probably coriander and dill. Many of these remains were found in the mineralised state confirming a human faecal context (Jacomet and Kreuz, 1999). The recorded species represent common elements Roman crop husbandry and arboriculture (Kuiper and Turner, 1992; White, 1970) and also occur frequently in the archaeobotanical finds in Roman to early Byzantine Sagalassos. Also, the presence of coprophilous fungi in the deep layers of the sewage channel provides clear evidence that the deposits are indeed associated with a faecal context (van Geel et al., 2003).

The high concentration of charred plant remains may result from the disposal of ashes into the channel. Alternatively, the char might have been discarded therein on purpose. Wood ash is a desiccant and raises the pH, which has the joint effect of reducing odours and destroying pathogens (Mehl et al., 2011). Similar practice was done in Roman kitchens where ashes from the stove were thrown into a cesspit to neutralise the smell (Scobie, 1986). Furthermore, the lowermost layers (150–190 cm depth) of the sewage channel (locus E) display a distinct increase in calcium concentration, which may indicate that lime was added. Lime has similar properties as wood ash and therefore, these observations could indicate that Sagalassians have used both ashes and lime to improve the operation and sanitary conditions of the public latrine.

4.6.2. The second phase: a manure producing unit

At a certain point in the history of Sagalassos, the function of public latrine was abandoned. It was shown by radiocarbon dating that this transition occurred in the early Byzantine period (see below). The latrine seats were removed and the original floor was covered with several layers of sediments. The stratigraphy of these layers was complex and contained a large amount of urban waste. Preliminary analyses suggested that the room was used as a collector of excrements, as some layers indicate the presence of both brushite and weddellite. While these minerals may be linked to a urinary origin, other sources or in situ mineralisation (McCobb et al., 2001, 2004) cannot be excluded. The present study, however, provides robust evidence of herbivore derived 5β-stanols in both the uppermost layers (0–30 cm depth) of the sewage channel (locus C) and the stratified heap (locus A). Values of the [5β:(5β + 5α)] stanol ratio around 0.5 were slightly lower than the sewage channel layers that contained human faeces, even though absolute concentrations of faecal biomarkers were higher (Table S1). These lower values were found to be inherently related to the herbivore origin of the faecal biomarkers (Evershed et al., 1997; Leeming et al., 1996). Indeed, values of the coprostanol:5β-stigmastanol ratio around 0.25 evidenced the presence of herbivore dung. Moreover, the predominance of epi-5β-stanol over 5β-stanols at locus A (Fig. 5c) provides evidence that the herbivore dung has been subject to intensive bacterial reworking, which could have occurred during the composting of the dung. A small contribution of porcine dung, however, cannot entirely be ruled out as bile acids were not detected (cf. Bull et al., 2002, see below). Yet, its proportion must have been rather limited since pigs only constituted between 20 and 30% of the livestock of Sagalassos during the Roman and Early Byzantine period (De Cupere, 2001).

The production of a fuel composed of dried animal dung can be ruled out since the well-constructed water supply to the room does not provide suitable drying conditions. Another hypothesis would be that the room was used as a stable. However, the complex stratigraphy containing different types of urban waste, the intensive microbial reworking of the ruminant dung and the addition of a calcium source do not support this functionality. In fact, such a deposit is more likely to be associated with a manure producing unit.
First of all, the scattering of potsherds in ancient off-site agricultural soils is considered as a marker for manuring. This theory was first proposed by Wilkinson (1982) and further strengthened by Bintliff and Snodgrass (1988). This theory entails the collection of animal and human excrements together with household rubbish, which was regularly spread across the cultivated landscape as fertiliser; and, after iterated application, produces a pattern of scattered off-site potsherd distributions. Bull et al. (2001) have validated this theory for manuring practices on Minoan agricultural terraces by the combined study of off-site potsherd scattering and faecal biomarker analysis. The present study further supports this theory by delivering the first evidence, based on faecal biomarker analyses, of the collection of excrements together with household waste in a manure production site. Indeed, the latrine deposits excavated above original floor level were associated with a large amount of urban waste, e.g. ceramics, animal bones, coins, glass, marble slabs, tesserae and metal artefacts. The fine ware ceramics found near the original floor displayed traces of water abrasion and surface trampling, suggesting that the material was collected elsewhere and dumped here. This was also observed in the sediments sampled for the current study, where many ceramic fragments were observed in the layers that contained the ruminant dung (Fig. 2, locus A and top 30 cm of locus C). However, further analyses are required to assess the application of such a mixed deposit to the farmlands of Sagalassos.

Secondly, the predominance of 3α-epimers in the herbivore dung layers (Fig. 5c) shows that the animal dung was subject to composting. This practice has already been observed in Minoan agriculture (Bull et al., 2001) and the importance of decomposition of organic matter to release its nutrients in arid or semi-arid cultural soils is considered as a marker for manuring. This theory entails the collection of excrements together with household waste in a manure production site. Indeed, the latrine deposits excavated above original floor level were associated with a large amount of urban waste, e.g. ceramics, animal bones, coins, glass, marble slabs, tesserae and metal artefacts. The fine ware ceramics found near the original floor displayed traces of water abrasion and surface trampling, suggesting that the material was collected elsewhere and dumped here. This was also observed in the sediments sampled for the current study, where many ceramic fragments were observed in the layers that contained the ruminant dung (Fig. 2, locus A and top 30 cm of locus C). However, further analyses are required to assess the application of such a mixed deposit to the farmlands of Sagalassos.

Finally, all the scatters of potsherds in ancient off-site agricultural soils is considered as a marker for manuring. This theory was first proposed by Wilkinson (1982) and further strengthened by Bintliff and Snodgrass (1988). This theory entails the collection of animal and human excrements together with household rubbish, which was regularly spread across the cultivated landscape as fertiliser and, after iterated application, produces a pattern of scattered off-site potsherd distributions. Bull et al. (2001) have validated this theory for manuring practices on Minoan agricultural terraces by the combined study of off-site potsherd scattering and faecal biomarker analysis. The present study further supports this theory by delivering the first evidence, based on faecal biomarker analyses, of the collection of excrements together with household waste in a manure production site. Indeed, the latrine deposits excavated above original floor level were associated with a large amount of urban waste, e.g. ceramics, animal bones, coins, glass, marble slabs, tesserae and metal artefacts. The fine ware ceramics found near the original floor displayed traces of water abrasion and surface trampling, suggesting that the material was collected elsewhere and dumped here. This was also observed in the sediments sampled for the current study, where many ceramic fragments were observed in the layers that contained the ruminant dung (Fig. 2, locus A and top 30 cm of locus C). However, further analyses are required to assess the application of such a mixed deposit to the farmlands of Sagalassos.

Apart from the primary objectives, further remarkable observations contribute to the understanding of lipid biomarkers in soils and their diagenesis. One of these observations is the apparent infiltration of 5β-stanols down the sewage channel, as demonstrated by the fact that the abundance of epi-5β-stigmastanol decreased with depth relative to epicoprostanol (Fig. 5d, loci C and E). This can most likely be explained by the infiltration of composted ruminant dung down the sewage channel, where it intermingled with human excrements. Similar observations were already made in a Roman cesspit from Stanwick where faecal signatures, based on 5β-stanols, were obtained underneath the cesspit (Bethell et al., 1994). Another remarkable fact is that bile acids were not detected in the present latrine deposits. This is a remarkable observation since bile acids are generally considered more recalcitrant than 5β-stanols (Eihmmani et al., 1997, 2000; Bull et al., 2003). A feasible explanation would be that they have completely leached out of the sediments. Whereas stanols are relatively insoluble, approximately 0.5–1.8 µg/l, the solubility of bile acids much higher. The solubility of the latter is furthermore increased in alkaline conditions due to their weak acid functionality (Table S3). The latrine deposits under study exhibit rather...
uniform pH values between 7.5 and 8.0. Bile acids are quite soluble in such a slightly alkaline environment. For instance, deoxycholic acid has a solubility of 2.7 g/l at pH 8.0 (Table S3). Therefore, the complete leaching of the bile acids would be a very reasonable explanation for the failure to detect them. This recognition poses a limitation on the use of bile acids as faecal biomarkers in neutral to alkaline soils which are subject to extensive water percolation. A similar mechanism might be valid for other compound classes bearing acid functionalities. The observed infiltration of stanols, however, seems to require another explanation since these compounds are far less soluble and thus less prone to leaching. But very little is known about the long-term effects of water percolation and environmental factors that influence leaching. Perhaps these molecules were transported by a downward physical migration of colloidal soil particles. This may be further promoted by action of soil fauna living in the upper soil horizons.

5. Conclusion

Faecal biomarker and archaeobotanical analyses were conducted to reconstruct the use and maintenance of the public latrine of the ‘Imperial Baths’ of Sagalassos. The presence of human derived 5β-stanols, such as coprostanol and epicoprostanol, in the sewage channel provided unambiguous evidence for human excrements. Radiocarbon dating shows that the latrine was already in use shortly after construction of the ‘Imperial Baths’ in 120–165 AD. Macroscopic plant remains and pollen, recovered from the sewage channels, represent a number of common species of the Roman diet in Sagalassos. Interestingly, deposits above the original floor of the latrine, dating to the early Byzantine period, contained clear evidence of ruminant dung, based on the presence of plant-derived 5β-stanols, such as 5β-stigmastanol and epi-5β-stigmastanol. The predominance of epi-5β-stanols over 5β-stanols in these layers further indicates that the animal dung has been subject to intensive bacterial reworking. Therefore, it can be assumed that the former latrine was reused as a manure production site. This hypothesis is further sustained by the indications for lime added to the compost heap as lime acts as a desiccant, increasing the pH and thus suppresses bad smell released from the digesting manure. The reuse of the latrine as a manure producing facility is congruent with the history of the city of Sagalassos. The presence of human faeces by means of biomarker detection. Environ. Int. 27, 647–654.

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Appendix. Supplementary material


