

Article

C_{14–22} n-Alkanes in Soil from the Freetown Layered Intrusion, Sierra Leone: Products of Pt Catalytic Breakdown of Natural Longer Chain n-Alkanes?

John F. W. Bowles ^{1,*}, Jessica H. Bowles ² and Andrew P. Gize ³¹ School of Earth and Environmental Sciences, University of Manchester, Manchester M13 9PL, UK² UF Chimie, Collège Sciences et Technologies, Université de Bordeaux, 33405 Bordeaux, France; Jessica.bowles@u-bordeaux.fr³ Consultant Geochemist, 3 Kingsway, Frodsham WA6 6RU, Cheshire, UK; andy_gize@msn.com

* Correspondence: john.bowles@manchester.ac.uk

Received: 26 January 2018; Accepted: 27 February 2018; Published: 6 March 2018

Abstract: Soil above a platinum-group element (PGE)-bearing horizon within the Freetown Layered Intrusion, Sierra Leone, contains anomalous concentrations of n-alkanes (C_nH_{2n+2}) in the range C₁₄ to C₂₂ not readily attributable to an algal or lacustrine origin. Longer chain n-alkanes (C₂₃ to C₃₁) in the soil were derived from the breakdown of leaf litter beneath the closed canopy humid tropical forest. Spontaneous breakdown of the longer chain n-alkanes to form C_{14–22} n-alkanes without biogenic or abiogenic catalysts is unlikely as the n-alkanes are stable. In the Freetown soil, the catalytic properties of the PGE (Pt in particular) may lower the temperature at which oxidation of the longer chain n-alkanes can occur. Reaction between these n-alkanes and Pt species, such as Pt²⁺(H₂O)₂(OH)₂ and Pt⁴⁺(H₂O)₂(OH)₄ can bend and twist the alkanes, and significantly lower the Heat of Formation. Microbial catalysis is a possibility. Since a direct organic geochemical source of the lighter n-alkanes has not yet been identified, this paper explores the theoretical potential for abiogenic Pt species catalysis as a mechanism of breakdown of the longer n-alkanes to form C_{14–22} alkanes. This novel mechanism could offer additional evidence for the presence of the PGE in solution, as predicted by soil geochemistry.

Keywords: n-alkanes; platinum; catalyst; soil; humic acid; Freetown intrusion; Sierra Leone

1. Introduction

The Freetown Peninsula, Sierra Leone consists of a thick (7 km) sequence of layered gabbroic rocks. Platinum-group minerals (PGM) have been located in these rocks and have shown to be altered by superficial weathering [1,2]. Alluvial PGM have been recovered from the streams draining the intrusion and they are notably different in mineral assemblage, composition, and size from the PGM in the unaltered host rocks [3,4]. This difference has created debate concerning whether supergene processes could be responsible for the observed mineralogical differences. It has been suggested [5–7] that some of the PGM from the host rocks were altered during weathering and the platinum-group elements (PGE) products transported in solution to a regime of changed Eh and pH where growth of a new PGM suite occurred. Others consider this to be a “scientifically untenable hypothesis” [8]. There is textural evidence in the alluvial PGM, however, that is consistent with low-temperature growth and re-solution and re-growth. Aware of the abundant organic material and microbial fauna beneath the tropical rain forest cover of the intrusion and the ease with which the PGE react with organic materials [9,10], some studies [11–13] have examined the possible role of humic and fulvic acid in taking the PGE into solution at 25 °C. It is clearly possible for the PGE to enter solution under these conditions within a short time frame. Importantly, recent work has shown the ability of bacteria to play a part in the growth of PGM [14,15]. There is a problem

in developing an overall mechanism because there is evidence of different stages in the alteration and re-growth process, but the connections between these stages are circumstantial. A reviewer's comments on a recent paper describing the mineralogy, geochemistry and genesis of the alluvial Freetown PGM [4] can be paraphrased as: "OK we accept there is a question to be answered but if organic acids are involved how could that work?" This paper is a novel attempt to answer that question and to suggest an alternative mechanism to a conventional organic geochemical approach. The question to be asked is less "how does humic acid permit alteration of the PGM or transport of the PGE?", but rather "how is humic acid broken down by the presence of the PGE in soils?"

2. Humic Acids in Soils

Humic acids are products of the breakdown of dead organic matter derived from the biosphere (e.g., leaf litter). In the living cell, the organic molecular structures are enzymatically controlled, either as independent molecules (leaf waxes) or biopolymers (proteins, carbohydrates, and lipids). The molecules comprising biopolymers have a reactive functional group (e.g., acid, alcohol) through which they are bound to the biopolymer. Upon death of a cell, the biopolymers are broken down (hydrolyzed) in to smaller polymers and free molecules (e.g., protein → peptide → amino acid). The free molecules and biopolymer breakdown products are overwhelmingly (>99.9%) degraded by microbial processes, ultimately to CO₂ and H₂O [16].

The molecules comprising biopolymers have reactive functional groups that can recombine non-enzymatically to form unorganized polymers, which become geopolymers. Molecules without reactive functional groups can escape degradation by adsorption and clathration within the randomly structured geopolymers. Initially, relatively low molecular weight geopolymers are formed (fulvic acids) which become humic acids as random polymerization continues to increase the geopolymer molecular weight. Further polymerization yields humins (protokerogen), and ultimately kerogen, the quantitatively dominant geopolymer. As polymerization increases molecular weight, the geopolymers become more difficult to isolate and characterize. Fulvic and humic acids are both soluble in a sodium hydroxide solution, whereas humin and kerogen are insoluble. Fulvic acids are soluble in hydrochloric acid, whereas humic acids and humins are insoluble [16,17].

The transition from fulvic acid to humic acid and to kerogens is a continuum, with definition dependent only on solubility. The geopolymer however retain in part the molecular structures that are inherited from the living sources. To deduce the original biological sources, bulk methods such as carbon and nitrogen stable isotope analyses can be used. If the original molecular structures are sought, a common technique is to thermally fragment the polymer (pyrolysis), followed by quantitative identification of the individual molecules by gas chromatography-mass spectrometry (GC-MS) [18]. Pyrolysis can produce artifacts. For example, a C₁₆ straight chain hydrocarbon with a terminal acid or alcohol can become thermally broken adjacent to the functional group to yield the C₁₅ hydrocarbon in the final GC-MS analysis, as well as any free C₁₅. Among the many compounds detected from pyrolysis of humic acids is an important group, the *n* (normal) -alkanes, where *n* is the number of carbons. These are straight chains, with no branches or cyclics, and have the general formula C_{*n*}H_(2*n*+2), where *n* ranges from 1 to 100 or more and their melting points increase with *n*. The *n*-alkanes with low values of *n* form gases, such as methane (CH₄, i.e., *n* = 1). In the range *n* = 5 to 16 the alkanes are liquids at 20 °C (e.g., octane, C₈H₁₈, *n* = 8) and for higher values of *n* they are solid and waxy.

The carbon number, *n*, can be indicative of the original biological source. Aquatic plants (e.g., algae and phytoplankton) yield high concentrations of *n*-alkanes in the nC_{15–17} range. Sub-aerial plants synthesize high boiling point waxes (nC_{27–31}) as protection from desiccation by the sun and wind, to cover leaves and fruit. Odd-carbon numbered *n*-alkanes usually dominate [19]. The reason is that the cell polymerizes acetic acid (C₂H₄O₂) to synthesize fatty (carboxylic) acids, which containing multiples of two carbon atoms, have even carbon numbers. The cell then removes the terminal acid functional group (losing 1 carbon), resulting in the odd carbon numbered predominance in the *n*-alkanes.

It is normal alkanes with $n = 27, 29$, and 31 that are of particular interest here and they are abbreviated to nC_{27} , nC_{29} , and nC_{31} . In their simplest form the alkanes consist of a long straight chain of carbon atoms with the hydrogen atoms attached along the length. The waxy, non-water soluble alkanes form in the leaves of vegetation [20–22], especially in the leaves of broad-leaved trees [23]. They can be found to be preserved in the soil as their melting points range from 59.5 to 67.9 °C.

Histograms showing the distribution of n -alkanes in tree leaves and soils are shown in Figure 1. The samples chosen for this illustration deliberately come from a wide geographic area and are from moist tropical forests. In all cases, the alkane distribution consists of the nC_{27} , nC_{29} , nC_{31} alkanes to the virtual exclusion of lighter alkanes, especially those lighter than nC_{22} . The lighter alkanes (nC_{12} to nC_{19}) are indicative of derivation from algae in marine or lacustrine environments. This summary includes recent studies that correspond with the established view of this subject [24–27].

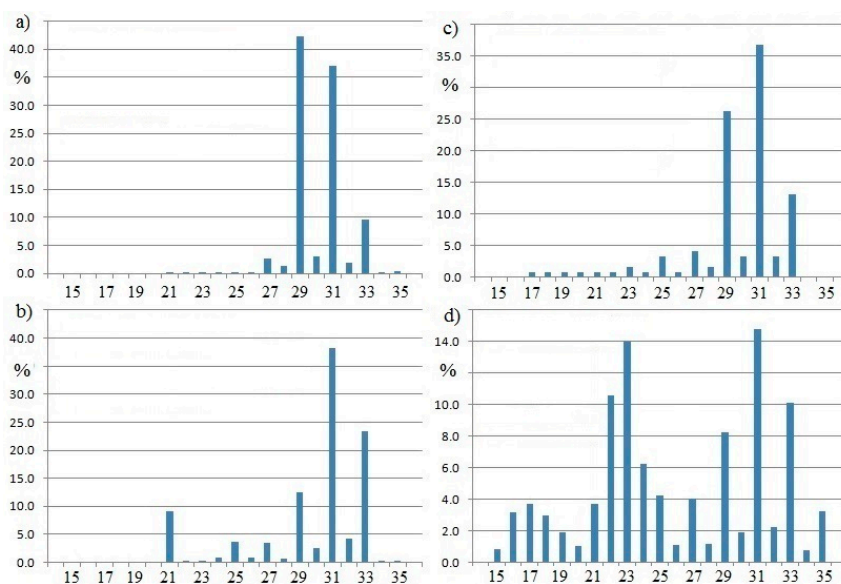


Figure 1. The typical percentage n -alkane content (nC_{14} to nC_{36}) of soils from moist tropical forests obtained using solvent extraction and analyzed by gas chromatography—mass spectrometry. (a) The average n -alkane content of 184 tree leaf samples from tropical lowland forest, Peru. Drawn from tabulated data [20]. (b) The average n -alkane content of leaves from seven tree species, Brazil. Drawn from tabulated data [21]. (c) The n -alkane content of a moist tropical forest near the coast at Debu, Cameroon. Drawn from tabulated data [22]. (d) The n -alkane content of five soil samples from moist tropical forest, Nepal. Drawn from tabulated data [23].

3. The Sample Site

The rocks beneath the sample site consist of a layered gabbroic sequence within which there is a PGE-enriched horizon containing small, weathered PGM [1,2,26]. The PGM are disseminated with rare minute Cu-sulfides. Pyrite and pentlandite are absent. Whole rock assays typically show the sulfur content below 0.04% with no known sulfide-rich layer. Weathering of these rocks does not produce a gossan. The rainfall is high (3000 to 5000 mm/year) [27], most falling in July to September. A thin soil, normally about 10 cm in thickness, overlies saprolite that is derived from weathering of the layered rocks. The humic and fulvic acid contents of a sample from that soil have been examined previously [12,13]. A comparison of the geochemistry of the rocks, saprolite, and soil has indicated that Pt, Pd and Au are mobile within the saprolite and soils [2]. The saprolite that covers the PGE-enriched horizon contains Pt, Pd, and Au. The saprolite retains details of the igneous layering and concentric weathering textures so it is in place and directly derived from the rocks that it covers. The saprolite that lies down slope of the PGE-enriched horizon also contains Pt, Pd and Au. This is not due to mechanical down slope movement, but it is the result of movement

of the metals in solution. There is an additional weathering feature; the upper levels of the saprolite contain less Pt, Pd, and Au than the lower levels and this is likely to have been caused by leaching under the heavy rainfall. The whole area is covered by closed canopy humid tropical forest [28]. The tree species mentioned in the older literature (*Khaya anthotheca*, *Guarea cedrata*, *Mimusops heckelii*, *Entandrophragma utile*, *Entandrophragma macrophyllum*, *Oldfieldia africana*, and *Cordia platythrsa*, [29] are mainly the hardwoods, such as mahogany and teak of commercial interest. The forest covering the sample area now consists mainly of secondary forest consisting of smaller trees with waxy, drip-tip leaves that are designed to cope with the high rainfall.

The sample site is on a well drained hilltop rising to 290 m above present sea level. The hill is a ridge (roughly N-S) which slopes steeply dropping by about 200 m on either side. This topography has existed for a long time. There are Eocene wave cut platforms and hard pan deposits at the base of the ridge (up to 50 m), and there is a lateritized raised beach at a height of about 200 m above present sea level. The sample site is at about 250 m above sea level and some 50 m above the lateritized raised beach (Figure 2). Field work is normally practical only outside of the rainy season, but at this time, the topography and freely draining saprolite mean that soil samples are dry. An attempt to sample ground water to determine the PGE contents and speciation was not successful.

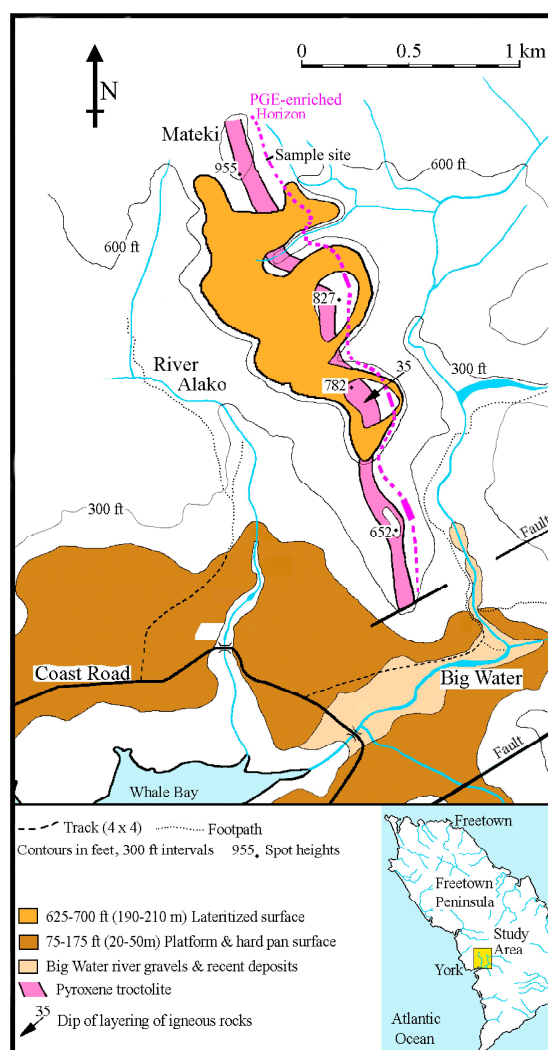


Figure 2. Map of the area inland from York on the Freetown Peninsula, Sierra Leone showing the sample site.

4. Methods

The soil sample was sieved to produce -6 mm and -177 μm (-80 mesh) fractions. Both of the fractions were assayed for Pt, Pd, and Au by Genalysis (Australia) using fire assay with a Pb collector using ICP-MS (Table 1). Assays of the reference material SARM 7 in the same run gave results that were close to the certified values.

Table 1. Soil sample assay (ppm).

Sample	Pt	Pd	Au
Sample 52, -6 mm to $+177$ μm fraction	52	6	1
Sample 52, -177 μm fraction	62	7	5

The humic acids were separated following the established method [30] and the dried humic acid analyzed by on-line pyrolysis gas chromatography-mass spectroscopy. The sample treatment and analysis have already been described [12,13].

The thermodynamic properties of some straight chain alkanes and their possible reactions with Pt-bearing species were studied by computational chemistry using the AMPAC 10 program [31]. This program offers several models corresponding to progressive development of the method. Two semi-empirical models (PM6 and RM1) from the AMPAC 10 program have been used here. Of these, PM6 is the most helpful for the present study because it can include calculations involving the PGE and Au, having been designed with biological molecules and catalysis in mind. There is a modification (D3H4) that is available to both of these models, which is intended improve the modelling of hydrogen bonding. Calculation of the Heat of Formation (Formation Enthalpy) of some n-alkanes from C_{11} to C_{32} indicates that the RM1 and PM6 models provide a good fit to values published previously [32], as shown in Figure 3. The D3H4 option does not offer an improvement in these examples. The Heat of Formation has previously been calculated [32] using a density functional model and when compared with experimental values derived from the NIST Chemistry Web Book (<http://webbook.nist.gov/chemistry>).

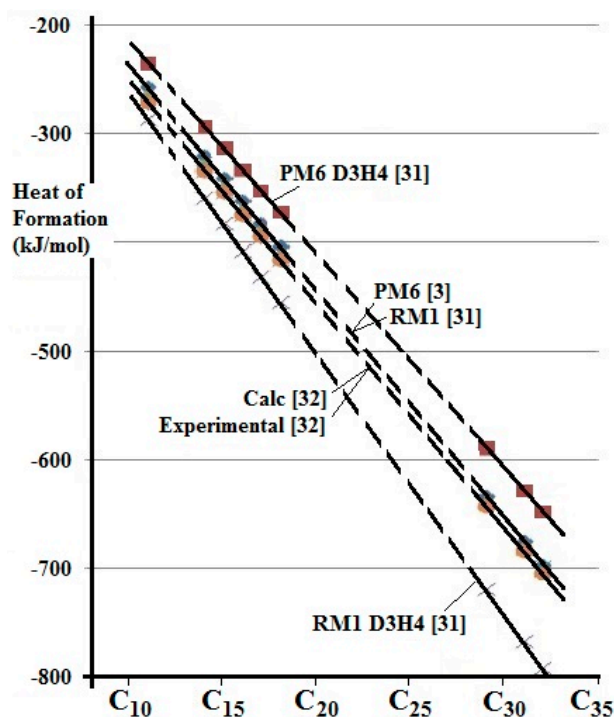


Figure 3. The variation of Heat of Formation for n-alkanes from C_{11} to C_{32} calculated using AMPAC 10 [31] and compared with earlier calculations and the experimental data [32].

5. Results

The Total Ion Chromatogram (TIC) trace for the humic acid from soil sample from the Freetown Peninsula, Sierra Leone, is reproduced as Figure 4a. The alkanes from the central trace in Figure 4a are represented as a histogram in Figure 4b for ease of comparison with Figure 1. nC_{27} , nC_{29} , and nC_{31} are present in larger proportions than their neighbours (nC_{28} , nC_{30} , nC_{32}), as is to be expected from the data in Figure 1. Surprisingly, the n -alkanes C_{14} to C_{24} are present in higher proportions than nC_{27} , nC_{29} , and nC_{31} , which is unusual. The high proportions of nC_{14} and nC_{15} are particularly noticeable given that their melting points are 5.9 and 9.9 °C, respectively. nC_{16} and nC_{17} with melting points of 18.2 and 21 °C are also present. As a measure for comparison with the n -alkanes in other broad leaved trees and tropical soils the ratio: average proportion of C_{14} – C_{26} /average proportion of C_{27} – C_{35} can be calculated from earlier results [20–23]. In the leaves of broad leaved trees, this ratio is 0.006 to 0.092 and it is 0.07 to 0.68 in tropical soils. For the sample from the Freetown Intrusion, this ratio is 3.34.

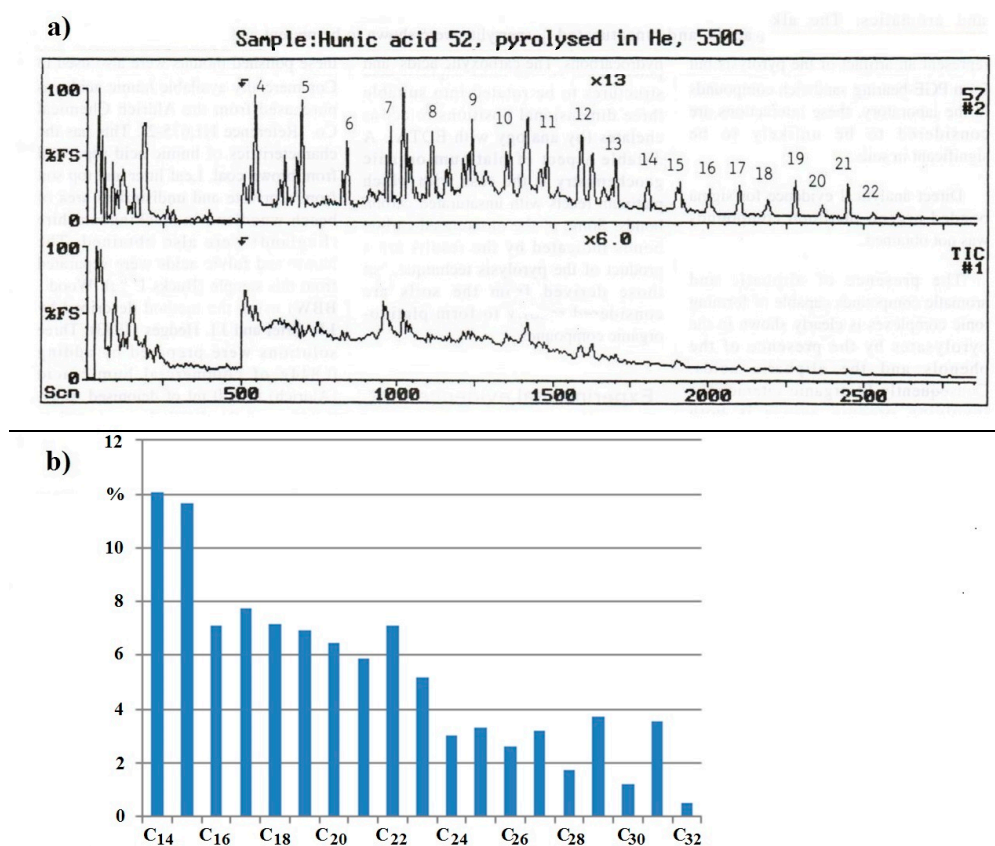


Figure 4. (a) Total ion current chromatogram (lower trace) and selected ion trace (upper trace) of a humic acid pyrolysate (sample 52). The lower trace shows the total ion current chromatogram with only a few peaks evident as a result of overlap from many compounds. A homologous series of normal alkanes is shown, starting from peak 4 ($\text{CH}_3(\text{CH}_2)_{12}\text{CH}_3$) to peak 22 ($\text{CH}_3(\text{CH}_2)_{30}\text{CH}_3$). The small shoulder to the left of some of the numbered peaks is the alkene with the same number of carbon atoms as the alkane. (b) The n -alkane (C_{14} to C_{32}) content of a soil sample from the Freetown Peninsula, Sierra Leone [12,13] presented as a histogram for comparison with Figure 1.

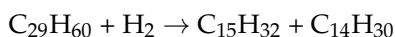
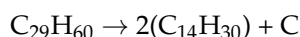
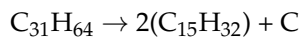
6. Discussion

6.1. Are nC_{14} – C_{25} Breakdown Products of nC_{27} – C_{31} ?

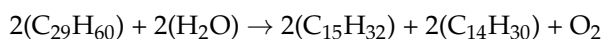
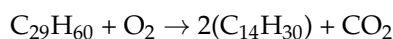
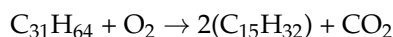
Tree leaves and their humic products normally contain the n -alkanes C_{23} – C_{31} and the range C_{27} – C_{31} is usually dominant. The C_{14} – C_{21} content of dry land vegetation is often negligible and

C₂₂–C₂₈ spasmodic in occurrence, depending upon the type of vegetation, but they are always lower in proportion when compared with C₂₇–C₃₁. The soil from the Freetown Peninsula is anomalous in that it contains three times as much C₁₄–C₁₅ as C₂₇–C₃₁, and about twice as much C₁₆–C₂₂ when compared with C₂₇–C₃₁. The sample site on a 200 m high ridge makes it unlikely that the presence of the lighter alkanes can be due to lacustrine algae. Although there are the remnants of raised beaches in the area these are at a lower level and of significant age in an area of very heavy rainfall so that lighter alkanes due to marine algae should not now be present.

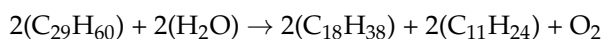
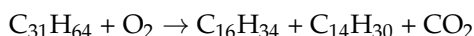
Could the presence of C₁₄ and C₁₅ be explained as a breakdown product of C₂₇–C₃₁ by reactions such as:



It would be more realistic to present these as oxidizing or reducing reactions:

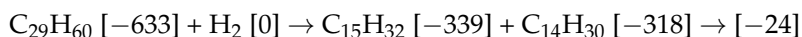
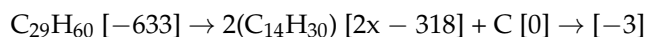
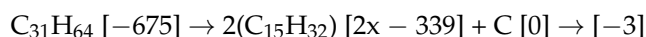


Similarly, the presence of C₁₆ and above might be explained in terms of a breakdown into two unequal parts, for example:

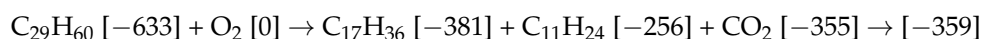
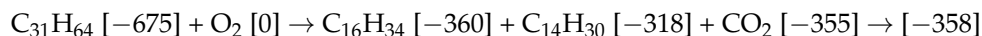
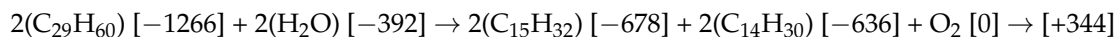
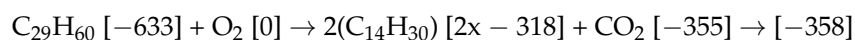
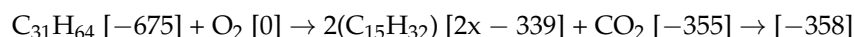


For this we would assume that the lighter alkanes <C₁₄ are sufficiently liquid to drain through the soil and that the alkanes C₁₄–C₁₇ are sufficiently viscous that they do not drain away or are being produced at a sufficient rate to compensate for any loss.

There is a huge problem with this suggestion. The alkanes, such as C₂₇–C₃₁, do not normally breakdown spontaneously because a large energy input is necessary to enable their destruction. Typically, this involves combustion. Calculation of the Heat of Formation of the components (Table 2) illustrates this problem. The reactions above can be shown with the calculated Heat of Formation (in square brackets and in kJ/mol).



For these reactions, there is an energy gain provided by breakdown but it is small.



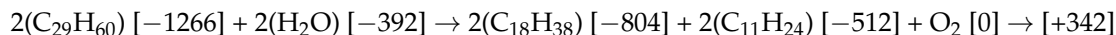
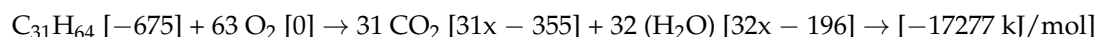


Table 2. Heat of formation (kJ/mol) calculated by AMPAC 10 [31] using the PM6, PM6 D3H4, RM1, and RM1 D3H4 models, and compared with calculated and experimental data [32].

Model	nC ₁₁	nC ₁₄	nC ₁₅	nC ₁₆	nC ₁₇	nC ₁₈	nC ₂₉	nC ₃₁	nC ₃₂
PM6	−255.60	−318.45	−339.38	−360.36	−381.29	−402.23	−632.67	−674.58	−695.51
PM6 D3H4	−233.67	−292.90	−312.67	−332.39	−352.15	−371.91	−589.20	−628.69	−648.45
RM1	−262.05	−324.78	−345.70	−366.60	−387.49	−408.42	−638.36	−680.19	−701.08
RM1 D3H4	−286.25	−358.44	−382.51	−406.58	−430.61	−454.69	−719.38	−719.38	−791.60
[28]	−270.68	−332.60	−353.07	−373.88	−394.44	−415.16	−642.59	−683.91	−704.60
[28]	−270.47	−332.35	−352.95	−373.55	−394.40	−414.83	−	−	−697.10

The oxidizing reactions with the addition of oxygen produce CO₂ and provide a heat gain. They are energetically favourable and the heat gain [−358 kJ/mol] is almost entirely due to the heat of formation of the CO₂. However, in practice, these reactions are only found to proceed if heat is added to trigger combustion. The reducing reactions are not energetically favourable and would require an additional +344 kJ/mol to proceed. The failure of these reactions to proceed without heat input explains why the non-oxidizing reactions with a smaller heat gain also fail to proceed. In these examples, different starting materials (C₂₉ or C₃₁) breakdown to equal or unequal components appears to make little difference. The oxidizing reactions above correspond only to partial combustion. Full combustion:



provides a very much greater heat gain.

The situation can be illustrated by the classic activation energy diagram (Figure 5), in which reactions proceed from left to right. The oxidation reaction leading to production of CO₂ is energetically more favourable but the reaction does not progress due to the energy requirements of the transition state. Only with the addition of heat (by combustion for example) can the transition state be reached. However, in the presence of a catalyst, the energy requirement of the transition state is reduced and the reaction could proceed without additional heat input.

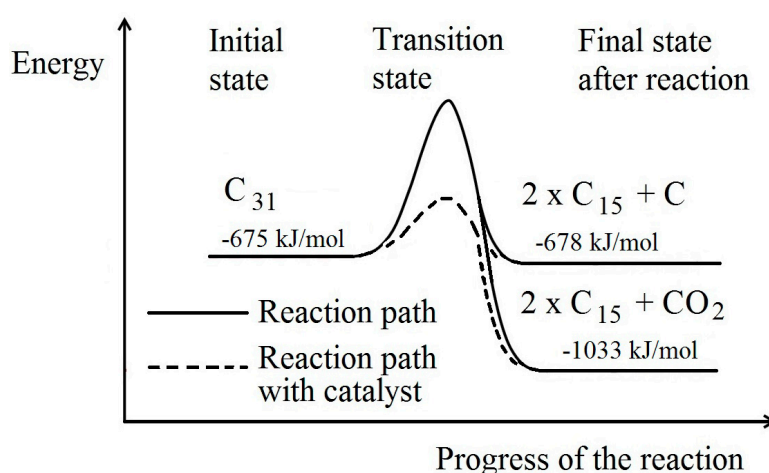
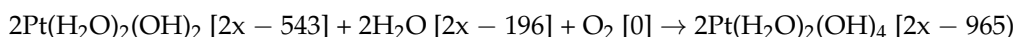


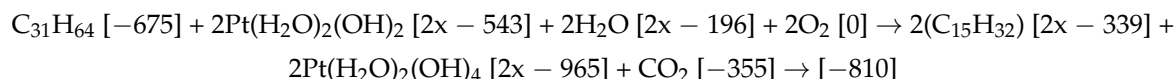
Figure 5. Schematic diagram illustrating the possible breakdown of C₃₁ to C₁₅ without oxidation and (lower line) with oxidation.

One distinguishing feature of the soil sample that is studied here is the Pt, Pd, and Au content. These elements are not likely to be present in significant amounts in most other soils, such as those

illustrated in Figure 1. Oxidized hydrous Pt species have a much lower heat of formation of than the un-oxidized species. For instance, [with Heat of Formation in kJ/mol and in square brackets]:



offers a gain of -452 kJ/mol. Thus, a linked Pt species-alkane reaction such:



has a much greater heat gain compared with the simpler alkane oxidation reactions described above. If such a reaction could occur, then it could provide the incentive for the breakdown of the alkane and overcome the energy of the transition state.

6.2. Can Catalysis Involving the PGE Facilitate Breakdown of the Heavier Alkanes?

Oxidation and ignition of four light alkanes in the presence of a Pt catalyst [33] has shown that the oxidation temperatures are lowered to: C_1 540°C , C_2 320°C , C_3 250°C , C_4 220°C , whereas these alkanes normally oxidize above 1000°C . The trend is expected continue so that the longer chain alkanes should oxidize at even lower temperatures. A straight line on a plot of temperature against C–H bond energy predicts a very low oxidation temperature for the longer chain alkanes [33]. The data suggest that a curve would be more appropriate, resulting in a gentler change of oxidation temperature with alkane number. A curve drawn through the data points by eye or using a Lagrange polynomial extrapolation suggests that C_{29} , C_{31} might oxidize at 195°C , but such an extrapolation is highly speculative and ignores both the large error bars on the data and the phase change from gas to solid. Somewhere between the straight line interpretation and a curve it appears likely that longer chain alkanes, such as C_{29} and C_{31} , could oxidize at room temperature in the presence of a Pt catalyst. Platinum was shown to be a more effective than Pd, Rh or Ir catalysts. At a slightly higher alkane/air mixture, the Pt catalyst oxidized C_2 at 265°C , whilst oxidation temperatures of 350°C (Pd), 410°C (Rh), and 450°C (Ir) were reported [33].

There are reactions available both in Nature and are used as commercial processes that involve Pt (or the other PGE) as catalysts. Some of these reactions are summarized below. A small quantity of PGE acting as a catalyst is not consumed, so it can alter a very much larger quantity of humic acid.

6.3. Industrial Catalytic Reactions Used to Modify the Alkanes

Hydrogenation breaks a carbon-carbon double bond using a catalyst, such as Pt, Pd, or Ni in the presence of excess H_2 . The organic molecule and the H_2 are adsorbed onto the catalyst surface and the H_2 dissociates, which allows the two hydrogen atoms to bond to different carbon atoms in the organic molecule, which is then released from the surface. Ni offers a slower reaction than Pt or Pd and requires a higher temperature. Industrial applications mainly involve the modification of the alkenes and there are variations that are adapted to create particular products, converting unsaturated compounds into saturated compounds (margarine, paraffin, naphthene), generally with better storage properties [34,35].

Dehydrogenation removes hydrogen from an organic molecule and is used to convert alkanes into more reactive products. Industrial processes use Ag, Fe + Mo or vanadium oxide as a catalyst, but require higher temperatures (250 – 400°C) to perform [36].

Hydrogenolysis breaks a single bond, typically a carbon-carbon single bond but C–O, C–N or C–S bonds can also be broken. Industrially, it is used to remove sulphur, but in the laboratory it is used for organic synthesis [37].

Alkane metathesis acts on a carbon-carbon single bonds and can be used both to generate a longer or a shorter alkane. A TaH (tantalum hydride) catalyst acts at 25 – 200°C and catalysts using either Pt in

conjunction with tungsten oxide or Ir with Rh are used in industry. In all cases, the presence of SiO_2 appears to be essential. The industrial processes are usually designed to produce longer chain alkanes, modifying C_9H_{20} to produce $\text{C}_{19}\text{H}_{40}$, for instance [38,39].

6.4. Possible Pt-Alkane Reactions in a Soil Environment

The physical situation envisaged in many of the industrial catalytic applications is a Pt foil with n-alkanes mobile in solution and the ends of the n-alkanes that are able to touch the Pt foil. In a humic-rich soil with the Pt mobile as a result of weathering, the Pt can be envisaged as free to move and able to interact with an n-alkane at any point along its length.

The AMPAC 10 program [31] has been used to construct the n-alkane molecules and calculate the Heats of Formation C_{11} to C_{32} listed in Table 2. The C_{31} molecule is shown in Figure 6a and it has a Heat of Formation of -674.54 kJ/mol .

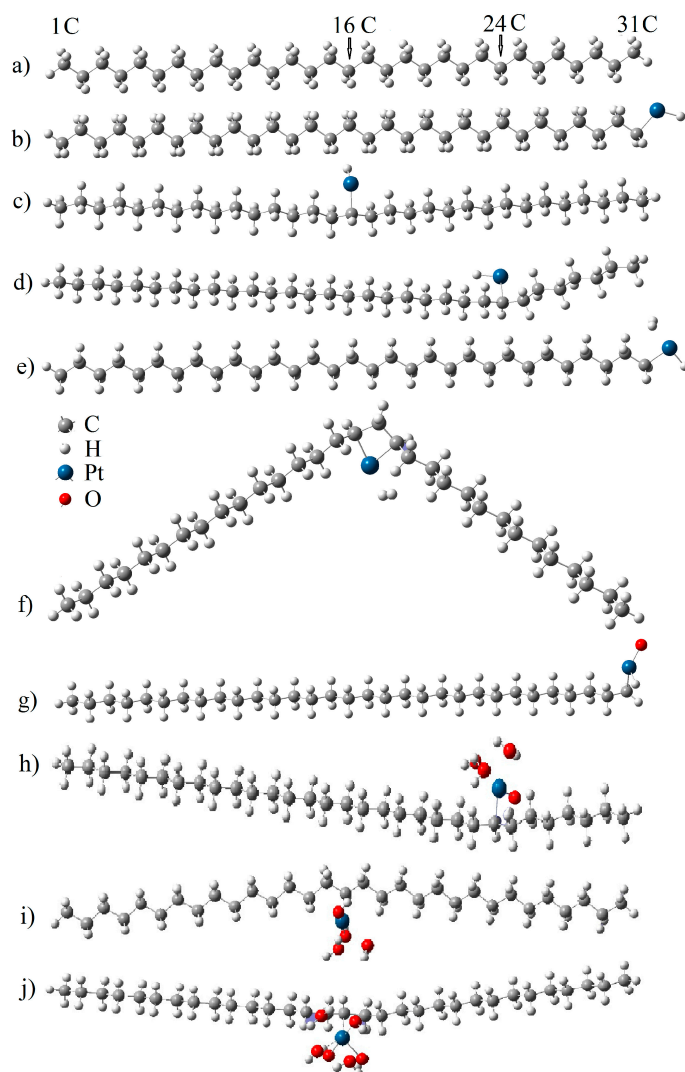


Figure 6. C_{31} molecules fabricated using the AMPAC 10 program [31]. The first, 16th (central), 24th ($\frac{3}{4}$) and the 31st carbon atoms are labelled 1C, 16C, 24C and 31C respectively. (a) $\text{C}_{31}\text{H}_{64}$, (b) $\text{C}_{31}\text{H}_{64}$ with Pt at end, attached to 31 C, (c) $\text{C}_{31}\text{H}_{64}$ with Pt at centre, attached to 16 C, (d) $\text{C}_{31}\text{H}_{64}$ with Pt at $\frac{3}{4}$, attached to 24 C, (e) $\text{C}_{31}\text{H}_{64}$ with tetrahedrally coordinated Pt at end attached to 31 C, (f) $\text{C}_{31}\text{H}_{64}$ with tetrahedrally coordinated Pt at centre attached to 16 C, (g) $\text{C}_{31}\text{H}_{64}$ with PtOH at end, attached to 31 C, (h) $\text{C}_{31}\text{H}_{64}$ with $\text{Pt}(\text{H}_2\text{O})_2(\text{OH})_2$ at $\frac{3}{4}$, attached to 24 C, (i) $\text{C}_{31}\text{H}_{64}$ with $\text{Pt}(\text{H}_2\text{O})_2(\text{OH})_2$ at centre, attached to 16 C, (j) $\text{C}_{31}\text{H}_{64}$ with $\text{Pt}(\text{H}_2\text{O})_2(\text{OH})_4$ at centre, attached to 16 C.

Allowing for a Pt atom to replace an H raises the Heat of Formation. If the Pt is added at the end (at the 31st C atom indicated here as 31C) the Heat of Formation becomes -347.80 kJ/mol (Figure 6b). At the centre (16C) of the alkane (Figure 6c), a Pt atom changes the Heat of Formation to -359.73 kJ/mol. In either case, there is little difference to the structure. If a Pt atom is added $\frac{3}{4}$ of the way along the alkane (24C) then the effect is to bend and twist the alkane (Figure 6d) and the Heat of Formation becomes -378.61 kJ/mol.

A tetrahedrally coordinated Pt atom at the end of the alkane (31C) makes little difference to the structure (Figure 6e, Heat of Formation -385.5 kJ/mol) but, in the centre of the alkane (16C), the alkane is both bent and twisted (Figure 6f, Heat of Formation -394.52 kJ/mol).

PtOH placed at the end of the alkane also has little effect on the structure (Figure 6g) with a similar Heat of Formation (-341.22 kJ/mol) to that given by Pt alone.

Platinum may occur in solution as $\text{Pt}^{2+}(\text{H}_2\text{O})_2(\text{OH})_2$ or $\text{Pt}^{4+}(\text{H}_2\text{O})_2(\text{OH})_4$. If $\text{Pt}^{2+}(\text{H}_2\text{O})_2(\text{OH})_2$ is placed at the $\frac{3}{4}$ position (24C, Figure 6h) or at the centre (31C, Figure 6i) of the C_{31} alkane the result is a bending and twisting of the alkane and the Heats of Formation are -1277.85 and -1294.56 kJ/mol respectively. $\text{Pt}^{2+}(\text{H}_2\text{O})_2(\text{OH})_4$ at the centre (Figure 6j) results in bending and twisting of the alkane and a significantly lower Heat of Formation (-1681.71 kJ/mol).

It seems that the addition of Pt species to the alkane can damage the structure, and that for Pt species, such as $\text{Pt}^{2+}(\text{H}_2\text{O})_2(\text{OH})_2$ or $\text{Pt}^{4+}(\text{H}_2\text{O})_2(\text{OH})_4$, the Heat of Formation is significantly lower. With such a low Heat of Formation, there is an increased likelihood of a reaction that is able to overcome the energy of the transition state and cause a breakdown of the n-alkane into smaller n-alkanes.

7. Conclusions

The soil above a PGE-bearing horizon within the Freetown Layered Intrusion contains high and anomalous concentrations of n-alkanes ($\text{C}_n\text{H}_{2n+2}$) that are in the range C_{14} to C_{22} . The presence of these n-alkanes cannot reasonably be attributed to an algal or lacustrine origin. Longer chain n-alkanes (C_{23} to C_{31}) in the soil are likely to have been derived from the breakdown of organic matter from leaf litter beneath the closed canopy humid tropical forest. These longer chain n-alkanes are typical of many other tropical to temperate soils.

Unassisted breakdown of the heavier n-alkanes to provide a source for the C_{14} to C_{22} alkanes is not likely. The alkanes are normally stable, and only breakdown when additional energy is added to the system, such as by ignition. Although there is an energy gain that is offered by breakdown, the stability of the heavier alkanes can be explained by the energy of a transition state, which is too high to be overcome without the input of additional energy.

In the Freetown soil, the catalytic properties of the PGE (and Pt in particular) may permit breakdown of the heavier alkanes by lowering the temperature at which oxidation can occur. Reaction between the heavier alkanes and Pt species, such as $\text{Pt}^{2+}(\text{H}_2\text{O})_2(\text{OH})_2$ and $\text{Pt}^{4+}(\text{H}_2\text{O})_2(\text{OH})_4$, can bend and twist the alkanes and significantly lower the Heat of Formation. Together, these effects render the breakdown of the heavier alkanes into smaller components more likely. It is possible, therefore, to offer an explanation for the presence of the C_{14} to C_{22} alkanes as a breakdown product of the natural, longer chain n-alkanes through the catalytic action of Pt, and provide an indication that Pt is present in solution in the soils, as predicted by the geochemical evidence [2].

The objective of this paper is to offer a preliminary assessment of the potential effects of solid and aqueous PGE (specifically Pt) on humic acids. Our ab initio approach was stimulated by the difficulty of identifying potential sources of biogenic matter for the enhanced concentration of n-alkanes that are centred on nC_{15} . Although future detailed work might identify biogenic sources, the present study shows that the nC_{15} enhancement can be formed abiogenically. The same argument could be applied to methyl alkanes and alkenes in the source leaf waxes. Fragmentation adjacent to weak bonds in long chain leaf waxes may occur, but we show that $\text{Pt}^{2+}(\text{H}_2\text{O})_2(\text{OH})_2$ and $\text{Pt}^{4+}(\text{H}_2\text{O})_2(\text{OH})_4$ can promote scission at the central carbon in nC_{31} . Other Pt species may occur in natural systems. Evidence for the conditions under which inorganic ligands, such as $\text{Pt}(\text{OH})$, $\text{Pt}(\text{OH})_2$, and $\text{Pd}(\text{OH})_2$, and organic

siderophiles can occur have been provided by experimental studies [40–45]. The initial study reported here did not find $\text{Pt}(\text{OH})$ or $\text{Pt}(\text{OH})_2$ to have a strong effect on the structure of nC_{31} (e.g., Figure 6g). $\text{Pt}^{4+}(\text{H}_2\text{O})_2(\text{OH})_4$, appears to be more effective and was selected for the ab initio calculations on the basis of a low desorption energy, thermodynamic stability, and the ability to dissolve platinum nanoparticles [46], a situation compatible with the platinum assays for the Freetown studies.

The calculated Heats of Formation assume a linear nC_{31} structure. Given the ease of C-C rotation in any alkane, it is possible that a linear nC_{31} molecule only exists in crystalline forms, such as leaf waxes, and not in randomly structured humic acids. Consequently, steric energy changes due to C-C rotation have not been included in our modelling. Our ab initio approach indicates abiogenic catalysis of long chain n-alkanes is energetically viable and fits the analytical data, which was our initial aim. Future work will need to include:

- the relative importance of the abiogenic, sterile processes presented in this paper compared with the non-sterile biogenic mechanisms;
- the relative efficiency of a range of oxide and hydroxide species, including especially $\text{Pt}(\text{OH})$, and $\text{Pt}(\text{OH})_2$;
- a comparison of Pt oxide and hydroxide species with those of the other PGE, especially Pd;
- an examination of the role of position along the alkane on the degree of distortion of the molecule;
- a comparison of the distortion of nC_{31} with nC_{29} and nC_{27} ; and,
- the effect of alkane structures other than simple linear chains.

Acknowledgments: The AMPAC 10 program [31] was kindly made available by Institute of Molecular Sciences (ISM), Theoretical Chemistry group (THEO) at the University of Bordeaux. The citation for this program (AMPAC 10, 1992–2013) is intended by the program developers to demonstrate that it is the result of a cooperative effort of a large, international team over many years. The editors and two anonymous referees have clarified the content of this paper and improved the presentation.

Author Contributions: The three authors have jointly written this paper, J.H.B. and J.F.W.B. used AMPAC10. APG separated the humic acids and conducted the on-line pyrolysis gas chromatography-mass spectroscopy.

Conflicts of Interest: The authors declare no conflict of interest.

References and Note

- Bowles, J.F.W.; Prichard, H.M.; Suárez, S.; Fisher, P.C. The first report of platinum-group minerals in magnetite-bearing gabbro, Freetown Layered Complex, Sierra Leone: Occurrences and genesis. *Can. Mineral.* **2013**, *51*, 455–473. [\[CrossRef\]](#)
- Bowles, J.F.W.; Suárez, S.; Prichard, H.M.; Fisher, P.C. Weathering of PGE-sulfides and Pt-Fe alloys, in the Freetown Layered Complex, Sierra Leone. *Miner. Deposita* **2017**, *52*, 1127–1144. [\[CrossRef\]](#)
- Bowles, J.F.W. The distinctive suite of platinum-group minerals from Guma Water, Sierra Leone. *Bull. Minéral.* **1981**, *104*, 478–483.
- Bowles, J.F.W.; Suárez, S.; Prichard, H.M.; Fisher, P.C. The mineralogy, geochemistry and genesis of the alluvial platinum-group minerals of the Freetown Layered Complex, Sierra Leone. *Mineral. Mag.* **2018**, *82*. [\[CrossRef\]](#)
- Bowles, J.F.W. The development of platinum-group minerals in laterites. *Econ. Geol.* **1986**, *81*, 1278–1285. [\[CrossRef\]](#)
- Bowles, J.F.W. Further studies of the development of platinum-group minerals in the laterites of the Freetown Layered Complex, Sierra Leone. In *Geo-Platinum 87*; Prichard, H.M., Potts, P.J., Bowles, J.F.W., Cribb, S.J., Eds.; Elsevier Applied Science: London, UK, 1988; pp. 273–280. ISBN 1851661972.
- Bowles, J.F.W. The development of platinum-group minerals in laterites: Mineral morphology. *Chron. Rech. Min.* **1995**, *520*, 55–63. [\[CrossRef\]](#)
- Cabri, L.J. pers. comm. 2015.
- Zeise, W.C.I. Von der wirkung zwischen platinchlorid und alkohol, und den dabei entstehenden neuen substanzen. II. Gekohlenwasserstoffes chlorplatin-ammoniak. *Ann. Phys. Chem. (Poggendorff)* **1831**, *21*, 497–549. [\[CrossRef\]](#)

10. Wood, S.A. The interaction of dissolved platinum with fulvic acid and simple organic acid analogues in aqueous solutions. *Can. Mineral.* **1990**, *28*, 665–673.
11. Bowles, J.F.W.; Gize, A.P.; Cowden, A. The mobility of the platinum-group elements in the soils of the Freetown Peninsula, Sierra Leone. *Can. Mineral.* **1994**, *32*, 957–967.
12. Bowles, J.F.W.; Gize, A.P.; Vaughan, D.J.; Norris, S.J. The development of platinum-group minerals in laterites; an initial comparison of the organic and inorganic controls. *Trans. Inst. Min. Metall. Sect. B Appl. Earth Sci.* **1994**, *103*, B53–B56.
13. Bowles, J.F.W.; Gize, A.P.; Vaughan, D.J.; Norris, S.J. Organic controls on platinum-group element (PGE) solubility. *Chron. Rech. Min.* **1995**, *520*, 65–73.
14. Reith, F.; Zammit, C.M.; Shar, S.S.; Etschmann, B.; Bottrill, R.; Southam, G.; Ta, C.; Kilburn, M.; Oberthür, T.; Bail, A.S.; et al. Biological role in the transformation of platinum-group mineral grains. *Nat. Geosci.* **2016**, *9*, 294–299. [[CrossRef](#)]
15. Campbell, G.; MacLean, L.; Reith, F.; Brewe, D.; Gordon, R.A.; Southam, G. Immobilisation of platinum by *Cupriavidus metallidurans*. *Minerals* **2018**, *8*, 10. [[CrossRef](#)]
16. Tissot, B.P.; Welte, D.H. *Petroleum Formation and Occurrence*, 2nd ed.; Springer: Berlin/Heidelberg, Germany, 1984; 699p, ISBN 3642878156.
17. Durand, B. *Kerogen: Insoluble Organic Matter from Sedimentary Rocks*; Editions Technip: Paris, France, 1980; 519p, ISBN 978-2710803713.
18. Fogel, M.L.; Sprague, E.K.; Gize, A.P.; Frey, R.W. Diagenesis of organic matter in Georgia salt marshes. *Estuar. Coast. Shelf Sci.* **1989**, *28*, 211–230. [[CrossRef](#)]
19. Peters, K.E.; Walters, C.C.; Moldowan, J.M. Source- and age-related biomarker parameters. In *The Biomarker Guide: Biomarkers and Isotopes in Petroleum Systems and Earth History*, 2nd ed.; Cambridge University Press: Cambridge, UK, 2005; Chapter 13; Volume II, pp. 483–607. ISBN 0-521-78158-2.
20. Feakins, S.J.; Peters, T.; Wu, M.S.; Shenkin, A.; Salinas, N.; Girardin, C.A.J.; Bentley, L.P.; Blonder, B.; Enquist, B.J.; Martin, R.E.; et al. Production of leaf wax n-alkanes across a tropical forest elevation transect. *Org. Geochem.* **2016**, *100*, 89–100. [[CrossRef](#)]
21. Jia, Q.; Sun, Q.; Xie, M.; Shan, Y.; Ling, Y.; Zhu, Q.; Tian, M. Normal alkane distributions in soil samples along a Lhasa-Bharatpur transect. *Acta Geol. Sin.* **2016**, *90*, 738–748.
22. Schwab, V.F.; Garcin, Y.; Sachse, D.; Todou, G.; Séné, O.; Onana, J.-M.; Achoundong, G.; Gleixner, G. Effect of aridity on $\delta^{13}\text{C}$ and δD values of C_3 plant- and C_4 graminoid-derived leaf wax lipids from soils along an environmental gradient in Cameroon (Western Central Africa). *Org. Geochem.* **2015**, *78*, 99–109. [[CrossRef](#)]
23. Da Silva, R.M.; de Tasso Moreira Ribeiro, R.; de Souza, R.J.C.; de Oliveira, A.F.M.; da Silva, S.I.; Gallão, M.I. Cuticular n-alkane in leaves of seven Neotropical species of the family Lecythidaceae: A contribution to chemotaxonomy. *Acta Bot. Bras.* **2017**, *31*, 137–140. [[CrossRef](#)]
24. Eglinton, G.; Hamilton, R.J. Leaf Epicuticular Waxes. *Science* **1967**, *156*, 1322–1335. [[CrossRef](#)] [[PubMed](#)]
25. Gelpi, E.; Schneider, H.; Mann, J.; Oró, J. Hydrocarbons of geochemical significance in microscopic algae. *Phytochemistry* **1970**, *9*, 603–612. [[CrossRef](#)]
26. Bowles, J.F.W. A primary platinum occurrence in the Freetown layered intrusion, Sierra Leone. *Miner. Deposita* **2000**, *35*, 583–586. [[CrossRef](#)]
27. Hughes, R.H.; Hughes, J.S. *A Directory of African Wetlands*; IUCN: Gland, Switzerland; Cambridge, UK; UNEP: Nairobi, Kenya; WCMC: Cambridge, UK, 1992; 820p, ISBN 2-88032-949-3.
28. Munro, P.G. Deforestation: Constructing Problems and Solutions on Sierra Leone’s Freetown Peninsula. *J. Polit. Ecol.* **2009**, *16*, 104–122. [[CrossRef](#)]
29. Clark, J.I. Vegetation. In *Sierra Leone in Maps*; Clarke, J.I., Ed.; Hodder and Stoughton: London, UK, 1969; pp. 24–25. ISBN 0340090855.
30. Ertel, J.R.; Hedges, J.I. Chemical and spectroscopic properties of marine and terrestrial humic acids, melanoids and catechol-based synthetic polymers. In *Aquatic and Terrestrial Humic Materials*; Christman, R., Gjessing, E., Eds.; Ann Arbor Science: Ann Arbor, MI, USA, 1983; pp. 143–163. ISBN 0250405504.
31. AMPAC 10 Semichem, Inc. 12456 W 62nd Terrace—Suite D; 1992–2013; AMPAC 10 Semichem, Inc.: Shawnee, KS, USA, 2009.
32. Abu-Awwad, F.M. The Gas-Phase Heats of Formation of *n*-alkanes as a function of the electrostatic potential extrema on their molecular surfaces. *E-J. Chem.* **2004**, *1*, 81–86. [[CrossRef](#)]

33. Veser, G.; Ziauddin, M.; Schmidt, L.D. Ignition in alkane oxidation on noble-metal catalysts. *Catal. Today* **1999**, *47*, 219–228. [[CrossRef](#)]
34. Anderson, J.R.; Boudart, M. *Catalysis—Science and Technology*; Springer: Berlin/Heidelberg, Germany, 1996; Volume 11, ISBN-13: 978-3-642-64666-9.
35. Hudlický, M. *Reductions in Organic Chemistry*, 2nd ed.; American Chemical Society: Washington, DC, USA, 1996; ISBN 0-86980-893-1.
36. Findlater, M.; Choi, J.; Alan, S.; Goldman, A.S.; Brookhart, M. Alkane Dehydrogenation. In *Alkane C–H Activation by Single-Site Metal Catalysis*; Pérez, P.J., Ed.; Springer: Dordrecht, the Netherlands, 2012; pp. 113–141. ISBN 978-90-481-3697-1.
37. Connor, R.; Adkins, H. Hydrogenolysis of oxygenated organic compounds. *J. Am. Chem. Soc.* **1932**, *54*, 4678–4690. [[CrossRef](#)]
38. Basset, J.M.; Copéret, C.; Lefort, L.; Maunders, B.M.; Maury, O.; Le Roux, E.; Saggio, G.; Soignier, S.; Soulivong, D.; Sunley, G.J.; et al. Primary Products and Mechanistic Considerations in Alkane Metathesis. *J. Am. Chem. Soc.* **2005**, *127*, 8604–8605. [[CrossRef](#)] [[PubMed](#)]
39. Burnett, R.L.; Hughes, T.R. Mechanism and poisoning of the molecular redistribution reaction of alkanes with a dual-functional catalyst system. *J. Catal.* **1973**, *31*, 55–64. [[CrossRef](#)]
40. Wood, S.A. Experimental determination of the hydrolysis constants of Pt^{2+} and Pd^{2+} at 25 °C from the solubility of Pt and Pd in aqueous hydroxide solutions. *Geochim. Cosmochim. Acta* **1991**, *55*, 1759–1767. [[CrossRef](#)]
41. Colombo, C.; Oates, J.; Monhemius, A.J.; Plant, J.A. Complexation of platinum, palladium and rhodium with inorganic ligands in the environment. *Geochem. Explor. Environ. Anal.* **2008**, *8*, 1–11. [[CrossRef](#)]
42. Azaroual, M.; Romand, B.; Freyssinet, P.; Disnar, J.-R. Solubility of platinum in aqueous solutions at 25 °C and pHs 4 to 10 under oxidizing conditions. *Geochim. Cosmochim. Acta* **2001**, *65*, 4453–4466. [[CrossRef](#)]
43. Wood, S.A.; van Middlesworth, J. The influence of acetate and oxalate as simple organic ligands on the behavior of palladium in surface environments. *Can. Mineral.* **2004**, *42*, 411–421. [[CrossRef](#)]
44. Dahlheimer, S.R.; Neal, O.; Fein, J.B. Potential mobilization of platinum-group elements by siderophores in surface environments. *Environ. Sci. Technol.* **2007**, *41*, 870–875. [[CrossRef](#)] [[PubMed](#)]
45. Gammons, C.H. Experimental investigations of the hydrothermal geochemistry of platinum and palladium: V. Equilibria between platinum metal, Pt(II), and Pt(IV) chloride complexes at 25 to 300 °C. *Geochim. Cosmochim. Acta* **1996**, *60*, 1683–1694. [[CrossRef](#)]
46. Koyama, M.; Kohno, H.; Ogura, T.; Ishimoto, T. Applications of computational chemistry to designing materials and microstructures in fuel cell technologies. *J. Comput. Chem.* **2013**, *12*, 1–7.



© 2018 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (<http://creativecommons.org/licenses/by/4.0/>).