Methanogenesis Potentials: Insights from Mineralogical Diagenesis, SEM and FTIR Features of the Permian Mikambeni Shale of the Tuli Basin, Limpopo Province of South Africa

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Abstract: Carbonaceous shale is more topical than ever before due to the associated unconventional resources of methane. The use of FTIR, SEM-EDX, and mineralogical analyses has demonstrated a promising approach to assess methanogenesis potentials in a more rapid and reliable manner for preliminary prospecting. Representative core samples from the borehole that penetrated the carbonaceous Mikambeni shale Formations were investigated for methanogenesis potentials. The absorption band stretches from 1650 cm$^{-1}$ to 1220 cm$^{-1}$ in wavenumber, corresponding to C-O stretching and OH deformation of acetic and phenolic groups in all studied samples, thereby suggesting biogenic methanogenesis. The CO$_2$ was produced by decarboxylation of organic matter around 2000 cm$^{-1}$ and 2300 cm$^{-1}$ and served as a source of the carboxylic acid that dissolved the feldspar. This dissolution process tended to release K$^+$ ions, which facilitated the illitization of the smectite minerals. The SEM-EDX spectroscopy depicted a polyframboidal pyrite structure, which indicated a sulfate reduction of pyrite minerals resulting from microbial activities in an anoxic milieu and causes an increase in alkalinity medium that favors precipitation of dolomite in the presence of Ca and Mg as burial depth increases. The contact diagenesis from the proximity of Sagole geothermal spring via Tshipise fault is suggested to have enhanced the transformation of smectite to chlorite via a mixed layer corrensite in a solid-state gradual replacement reaction pathway. The presence of diagenetic chlorite mineral is characteristic of low-grade metamorphism or high diagenetic zone at a temperature around 200 °C to 230 °C and corresponds to thermal breakdown of kerogen to methane at strong absorption band around 2850 cm$^{-1}$ and 3000 cm$^{-1}$, indicating thermal methanogenesis.

Keywords: methanogenesis potentials; biogenic methanogenesis; thermal methanogenesis; carbonaceous shale; carboxylic; illitization; chloritization; framboidal pyrite

1. Introduction

Carbonaceous shale has been more topical than ever before due to the associated unconventional resources for energy supply. Shale constitutes an essential component of unconventional gas generation due to the global clamor for an energy transition from heavy
greenhouse domination to low-carbon natural gases [1–3]. With respect to climate change moderation, carbonaceous shale has been identified to promote a short-term accelerated action to achieve the 2015 Paris agreement climate targets of keeping the global temperature increases below 2 °C. In recent studies [4,5], shale gas exploration has been prioritized in the US from 2006 to 2020, cutting off well over 430 million tonnes of CO₂ emission and accounting for over 30% of the financial fiscal year [6,7]. In contrast to conventional shale source rocks, carbonaceous shales tend to host natural gas under a low pressure and extremely low permeability but extractable using horizontal well-drilling and hydraulic fracturing technologies [8–11].

Despite many decades of research in carbonaceous shale, a reliable and rapid discriminative approach to identify a potential methane gas-bearing rock remains elusive. However, some studies have used petrographic and sedimentological properties [12,13], geophysical properties [14–16], and geochemical and mineralogical characteristics [17–20] to attribute most organic matter transformation to their burial. Furthermore, there is extensive recent literature [21–29] on the successful use of an organic geochemical method to determine petrophysical properties such as total organic carbon (TOC), thermal maturity, vitrinite content, and desorbed gas composition. Undoubtedly, these investigative approaches provide deterministic information rather than discrimination at the preliminary exploratory stage [30]. The discriminative approach identifies sweet-spot targets for further detailed and determinative investigations, which require huge cost implications.

This research provides information on methane formation during the transformation of organic components of organic-rich sedimentary deposits. It applies evidence from mineralogical, structural, and functional-group features of the carbonaceous Mikambeni shale in the Tuli Basin. The Permian Tuli Basin is known as the Limpopo-area Karoo Basin, hosting coal resources used to power a coal-fired plant for electricity generation in the northeastern region of South Africa [31–34]. The potential for methanogenesis sources has not been given proper attention in the burial depth, having a thickness of 1 km or less.

Study Area

The Tuli Basin is a trans-frontier depozone that straddles the triple junction of South Africa, Zimbabwe, and Botswana situated within latitudes 29°36′22.19″ E and 28°25′23.15″ E and longitudes 23°11′24.45″ S and 22°43′24.54″ S (Figure 1). This basin is a back-bulge structural depression that resulted from crustal subsidence due to tectonic regimes adjacent to the Cape Fold Belt (CFB) orogenic belt. The subsidence controls the accommodation space that preserves the Carboniferous–Jurassic sedimentary sequence of the Tuli Basin in South Africa [35,36]. The succession of formations from oldest to youngest is the Tshidzi, Madzaringwe, Mikambeni, Fripp, Solitude, Klopperfontein, Bosbokpoort, and Clarens Formations [37] (Figure 1).

The Mikambeni Formation consists of the Permian black carbonaceous shale intercalated with coal seams and alternating sandstone and siltstone [31]. The thickness of the Mikambeni Formation ranges from 60 to 100 m, corresponding to the proximal and distal points. The variation in shale thickness is interpreted to be caused by basin tectonics and differential subsidence trends as transporting river flowed from the northwest to deposit more sediments towards the southeast direction of the basin [34]. The coal seam associated with the Tuli Basin is dominated by colliotellinite, suggesting organic matter input from cellulose and woody tissues of the higher terrestrial plant. Based on lithofacies description, the shale color and content suggest a reducing condition associated with oxygen deficiency causing decay of fauna and flora or plant organisms. Given the overall sedimentary sequence, the deposition represents a fluvial-to-aeolian milieu. This deposition reflects different paleo-climatic overprints of different ages spanning from the Permian to the Jurassic Period [38]. Besides the lignin, the cellulose and tannins of vascular plants *dicroidium* are partly responsible for the main coal seam associated with the carbonaceous shale of the Mikambeni Formation [31,39]. In contrast to lithofacies of Mkambeni, the Tshidzi Formation shows non-sorted to poorly sorted argillaceous or sandy
sediments with particles size up to 2 m in diameter. This argillaceous diamicite suggests a glacial or fluvio-glacial environment resulting from a low energy level of mudflow to deposit well-sorted sediments. There is a progressive change from reducing or anoxic to the oxidizing environment as shale formations become younger. The Triassic black shale at the lower unit of Solitude Formation has a dark grey shale and grey shale towards overlying Bosbokpoort and Clarens Formations [38]. This color variation reflects a progressive fall in water level where sediments are deposited on floodplains of meandering rivers under dry and oxidizing conditions. The preponderance of red calcareous concretions of overlying the Bosbokpoort further support an oxidizing environment towards the upper Tshipise Basin.

![Figure 1. Map and sedimentary sequence of the Tuli Backbulge Basin.](image)

2. Materials and Methods

Mineralogical identification of 16 core samples from the borehole that penetrated the carbonaceous Mikambeni shale Formations at different depths was determined using X-ray diffraction (XRD) at Council for Scientific and Industrial Research (CSIR) Pretoria (Figure 2).

Based on the lithological facies, 8 representative shale samples corresponding to the formations were selected and pulverized in an agate mortar. The clay fractions were dispersed in 0.7 mg/mL of distilled water and put in an ultrasonic water bath for 40 s to prevent the flocculation of particles [40]. Afterward, the air-dried samples were treated with ethylene glycolation to identify smectite minerals and dimethyl sulphoxide to differentiate kaolinite from chlorite [41]. Samples were tightly mounted on oriented sample holder with very little pressure using black loading preparation technique using PANalytical X’Pert Pro-powder diffractometer equipped with X’Celerator detector coupled with receiving slits, variable divergence, and Fe-filtered Cu-Kα radiation at the Energy Centre CSIR Pretoria. While the receiving slit was positioned at 0.040°, the counting area was from 2 to 80° on a 2θ scale at 1.5 s [42]. The interstratified I/S minerals were measured on a less than 2 µm fraction of air-dried samples using Cu-Kα radiation [43]. Semi-quantitative
mineral measurements were obtained using single line fitting of TOPAS software while the detection limits remained 1%.

![Stratigraphic column of the studied borehole showing lithologic description and carbonaceous shale sampling depth.](image)

The functional group information was provided by the Fourier Transform Infrared (FTIR) Alpha Bruker spectrophotometer at the University of Venda, Limpopo. Pulverized samples were placed in the measurement position of the spectrometer and scanned between a minimum and maximum wavelength range of 500 and 4000 cm$^{-1}$, respectively. The absorbance spectrum depicts the crystalline phases which reflect their functional group.

A Zeiss EVO MA15 scanning electron microscope (SEM) equipped with a tungsten filament and a Bruker energy-dispersive X-ray (EDX) spectrometer were used for the morphological analysis. The system was operated under a high system vacuum, approximately 7.03 e to 007 mbar. The electron beam was generated with an accelerating voltage of 20 kV and a probe current of approximately 2nA. The sample was sprinkled onto a carbon tape substrate and carbon-coated prior to being loaded into the SEM. A secondary electron detector was used to produce images at 300×, 600×, and 900× magnification. Some of these particles are out of focus, owing to the three-dimensional depth loss of informa-
tion; focusing one part results in the other being out of focus on some of these images, particularly at high magnifications.

3. Results

The mineralogical identification of the carbonaceous shale samples obtained from different depths of two boreholes analyzed using X-ray diffraction (XRD) values and patterns is presented in Table 1 and Figure 3a,b. The results show the presence of clay minerals composed of montmorillonite, mixed illite/smectite (I/S), illite, chlorite, and non-clay minerals such as dolomite, albite, microcline, pyrite, and quartz. The average quartz content was 29.42%, ranging from 25.10% to 35.60%. However, quartz with higher values was found at a depth of 470 m. The relatively high quartz content may be attributed to the preserved detrital silica input from associated siltstone facies, since it has high resistance to mechanical alteration, although diagenetic quartz can precipitate from the dissolution of feldspar [38,44].

![Figure 3](image-url)

**Figure 3.** Cont.
In contrast to the quartz content, the average amount of microcline and albite was 10.17% and 5.45%. However, there is no albite mineral present at a depth of 470 m. Feldspars are susceptible to dissolution reaction as burial depth increases, as the albite minerals progressively reduce until the 470 m [45,46]. As burial depth increases, the montmorillonite content progressively decreases from 21.35% to 5.40%, while, on the other hand, the illite and mixed I/S contents increased from 15.20% to 20.50% and 8.25 to 13.20%. This observation is ascribed to the illitization reaction at burial depth temperature less than 200 °C in which precursor minerals of K-feldspar or smectite transform to illite or
mixed Smectite/illite (S/I) in the presence of sufficient K⁺ [47–49]. The chlorite content has an average value of 3.42% and ranges from 2.20% to 5.65% in a dispersed sequence in all samples. The presence of chlorite mineral suggests a by-product of the montmorillonite-illitization-chloritization transformation. This transformation process occurs in a closed system having a low ratio of fluid to shale rock [50]. The distribution of dolomite showed a progressive increment in concentration from 5.30 to 11.05 with burial depth. Since the organic-rich shale has a low fluid-to-rock ratio, the idea of dolomite replacing calcite with the saturated fluid of Mg²⁺ may not be feasible. The explanation is the precipitation of primary dolomite and stepwise recrystallization due to bacterial mediation [51]. It is worth noting that a high fluid–rock ratio is necessary for a steady and saturated supply of Mg²⁺ for dolomitization replacement of calcite, which favors permeable sandstone over impermeable buried shales. High burial temperature is insufficient to destroy the evidence of pyrite grain history or origin. Hence, its presence suggested detrital grain devoid of oxidizing setting but associated with anoxic, aridity, or fluvial settings [52,53].

The Fourier Transform Infrared (FTIR) spectra of the studied carbonaceous shale depict different absorbance bands of organic and mineral content at the various diagnostic wavelength of the infrared spectrum (Figure 4). Each absorbance band indicates the functional group representing the whole molecular structure of a material [30]. Most of the studied samples exhibit a strong characteristic absorbance peak around 650–900 cm⁻¹, corresponding to the aromatic phenolic group, C=C-OH. However, the peak reduces as temperature increases with depth due to bond deformation [54]. The aromatic C=C absorption group suggests a wavelength band that is typical of terrigenous plants that have lignin-derivative materials [55]. The peak stretching from around 1000 cm⁻¹ to 1220 cm⁻¹ indicates the C-O stretching and OH deformation of the carboxylic, COOH groups, suggesting decarboxylation of organic matter [56]. The deformation of the carboxylic group releases the CO₂, reflecting a characteristic strong absorption around 2000–2300 cm⁻¹ [30]. However, the aliphatic hydrocarbon is considered to contain more carboxylic- and phenolic-OH functional groups with a micelle-like structure that has a relative macromolecular weight ranging from 3500 to 100,000 Da compared with aromatic hydrocarbon [57]. In aromatic hydrocarbon, the voids among the carbon chain serve to act as trapping and binding sites for organic lipids and inorganic materials such as hydrous oxides and clay minerals [57]. The hydrocarbon gas, CH₄, shows infrared absorbance peaks between 2850 and 3000 cm⁻¹ due to aliphatic C-H stretching of methyl and methylene vibration [54,58]. The low absorption peak around 3500 cm⁻¹ corresponds to water desorption wavelength [59]. The spectra showed inconspicuous peaks of the carboxylic group at greater burial depths around 450 m, suggesting over-maturation of organic matter beyond the methanogenesis window.

Representative SEM images of the studied carbonaceous shales are presented in Figure 5a–d. The image depicts a mixture of pore structures, organic matter, and mineral contents. The image (Figure 5a) revealed conspicuous intragranular pores among grains of quartz, feldspars, chlorite, illite, carbonate, and pyrite minerals as shown by the XRD, forming an interconnected pore. The interconnected pores provide migration pathways for gaseous hydrocarbon during organic matter (OM) maturation [60]. The carbonate compound of calcite and dolomite was found to dominate the shale, with carbon elemental content ranging from 65% to 73% (Figure 6c). Furthermore, the organic matters were also hosted in the intragranular pores of mineral grain, which suggests dissolution cavity of carbonate minerals since intrapore of clay minerals such as feldspar is difficult to observe [61]. Non-spherical, closely packed polyframboidal pyrite was observed, which is affirmed by the multiple occurrences of Fe-elemental contents on the EDX result. The presence of the polyframboidal pyrites suggests an anaerobic condition that favors organic matter degradation and thermal maturation as burial depth increases [62]. The thermal maturity of the organic matter is further supported by the presence of fibrous chlorite and illite minerals (Figure 6d), which are index minerals for high-diagenetic temperature zone and quality hydrocarbon reservoirs [63].
Figure 4. FTIR spectra of the studied carbonaceous shale showing functional groups representing molecules (R-OH-phenolic, C-C-OH-acetic, carboxylic, carbon (IV), COOH-carboxylic, CO₂-carbon (IV) oxide, CH₄-methane, H₂O-water).

Figure 5. SEM images of the studied carbonaceous shale showing (a) intergranular pores hosting OM, (b) intragranular pores hosting OM, (c) SEM images of Chlorite-chl, Quartz-Q, Illite-I minerals, and (d) veinlet of Illite minerals.
Figure 6. (a) vertical, cylindrical pipes with open apices, (b) simple, mostly vertical shaft-like trace fossils, (c) close up of the wall of the digit-like structure showing the extensively branching burrow system with random orientation (d) Large-scale trace fossils showing lateral bifurcations on a sub-horizontal surface (modified from [37]).

4. Discussion
4.1. Diagenetic Alteration Pathways

The mechanical compaction of detrital sediments and terrestrial plant materials following sedimentation of the 40 m thick Fripp Formation over Mikambeni Formation initiate their diagenetic changes. At this initial stage, the brackish interstitial fluid or pore water and bacterial activities are pronounced due to the degradation of plant material to produce humic material. Humic substances are composed of weak acidic electrolytes with aliphatic and aromatic hydrocarbon, which may incorporate atomic radicals N, O, or S [57]. The humic acid gains significance at a wavenumber absorption peak around 650–900 cm\(^{-1}\), corresponding to the acetic and phenolic group. The compaction process leads to dehydroxylation in which the pore fluid is released, as indicated by the water absorbance band around 1300–1900 cm\(^{-1}\). More evidence of the early-stage organic degradation in the Tuli Basin was marked by bioturbation of burrowing organisms (Figure 6). In the studied basin, soft sediment burrowing organisms such as beetles and ants have produced un-bifurcated ichnofossils which are tentatively recognized as *Skolithos* sp. and other bifurcated trace fossils [36].

With an increase in burial pressure and depth of about 360 m thick, which results from successive overlying sediments of the Solitude, Klopperfontein, Bosbokpoort, and Clarens formations, the elevated temperature causes further geochemical and mineralogical changes. The gradual decrease in montmorillonite and a parallel progressive increase of illite minerals were observed, while mixed-layered montmorillonite-illite (M-I) mark the extent of progress along with series of reactions between burial depths of about 370 m to 470 m. In Mikambeni shale, the first series of the illitization reaction progress is made possible due to the presence of microcline and montmorillonite minerals as expressed in Equation (1) [64]. Alternatively, diagenetic illite may have crystallized from geochemical interaction between kaolinite and microcline or potassium ion (K\(^+\)), but the absence of
kaolinite inhibits that reaction’s progress pathway. Retrospective studies [44,65] have
reported the kaolinite illitization via reactions with k-feldspar, K⁺, or H⁺ as burial depth
increases; however, this reaction mechanism is hindered due to the absence of kaolinite
minerals in Mikambeni shale. Kaolinites may have precipitated due to increasing quartz
concentrations, but it is likely to be consumed immediately after crystallization as expressed
in Equation (2) due to high temperature. Studies [66–68] have unequivocally demonstrated
that the kaolinite dissolution in shale rock can take place at a lower temperature than 60 °C.

\[
\text{Al}_2\text{Si}_4\text{O}_{10}[\text{OH}]_2 + \text{H}_2\text{O} + \text{KAlSi}_3\text{O}_8 \rightarrow \text{KAl}_2\text{Si}_5[\text{OH}]_2 + \text{SiO}_2 + \text{H}_2\text{O} \quad (1)
\]

\[
\text{Al}^{3+} + \text{SiO}_2 + 2.5\text{H}_2\text{O} \rightarrow 0.5\text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4(\text{kaolinite}) + 3\text{H}^+ \quad (2)
\]

Furthermore, the organic and inorganic interaction promotes the illitization by initi-
tiating the release of the H⁺ ion from the carboxylic acid functional group as expressed
in Equation (3). The carboxylic C=O band can remain approximately at a wavenumber
of 1000 to 1225 cm⁻¹ stretching regardless of the source material. The CO₂ produced by
decarboxylation of organic matter is commonly suggested as a source of the acid that
dissolves the microcline feldspar [69]. This carbon dioxide, CO₂, is recognized from its
characteristic strong absorptions around 2000 and 2300 cm⁻¹.

\[
\text{CH}_2\text{O (carboxylic group) + H}_2\text{O} \rightarrow \text{CO}_2 + 4\text{H}^+ \quad (3)
\]

The presence of a carboxyl group increases solubility properties. As such, the labile
microcline mineral interacts with the pore water concentrated with carboxylic and phenolic
acids to produce silica and potassium cation, which enhances illitization of montmorillonite
mineral as represented in Equation (4). This resulting authigenic silica likely contributes to
the increasing quartz content with burial diagenesis, although the presence of preserved
detrital quartz inputs cannot be discounted. With the silstone facies interbedded with
the carbonaceous shale, the input of detrital silica is possible since quartz is largely stable
at high temperatures. Nonetheless, the increasing quartz contents tend to significantly
improve the shale reservoir quality because it imparts a brittleness property and enhances
the fracability of shale [70].

\[
\text{KAlSi}_3\text{O}_8 (\text{microcline}) + \text{H}^+ \rightarrow \frac{1}{2}\text{SiO}_2 + \text{K}^+ \quad (4)
\]

As the reaction continues, more K⁺ ions are concentrated via replacement K-feldspar
(microcline) by the albite at a higher temperature between 120 °C and 180 °C as represented
in Equation (5) [66,71]. The absence of kaolinite minerals with a parallel increase in chlorite
suggests complete dissolution of kaolinite at such a high temperature.

\[
\text{KAlSi}_3\text{O}_8 (\text{microcline}) + \text{Na}^+ \rightarrow \text{NaAlSi}_3\text{O}_8 (\text{Albite}) + \text{K}^+ \quad (5)
\]

With further burial diagenesis, the presence of a high concentration of potassium
K⁺ fluid promotes the illitization processes of montmorillonite minerals. However, other
factors such as time, fluid/rock ratio, nature of starting material, and kinetic mechanism
have been considered as important for illitization. It is worth noting that the direct
conversion of smectite to illite as expressed in Equation (6) [72] does not suggest a solid-state
reaction but rather shows the quantitative and compositional changes in shale mineralogy
that takes place during burial diagenesis. Importantly, this conversion of smectite to illite
reaction has been indicated to take place at similar temperature intervals as the matura-
tion index for organic matter and the release of interstitial water during I-M diagenesis
to enhance hydrocarbon migration [69]. By implication, the range for the change in I/S
ordering from random to regular during smectite illitization in shales coincides with the
temperature range of organic matter maturity and the onset of oil generation in young sedimentary basins.

\[
2\text{Na}_{0.4}(\text{Al}_{1.47}\text{Mg}_{0.50}\text{Fe}_{0.18})\text{Si}_{4}\text{O}_{10}(\text{OH})_{2}\text{(Smectite)} + 0.85\text{K}^{+} + 1.07\text{H}^{+} \\
\rightarrow 1.06\text{K}_{0.80}(\text{Al}_{1.98}\text{Mg}_{0.53})\text{Si}_{10}\text{O}_{25}(\text{OH})_{2}\text{(illite)} + 4.6\text{SiO}_{2} + 0.36\text{Fe(OH)}_{3} + 0.56\text{Mg}^{2+} + 0.8\text{Na}^{+} + 0.9\text{H}_{2}\text{O} \\
(6)
\]

Although present in low quantities (Table 1), chlorite mineral is increasingly crystallized as a discrete phase rather than evolutionary trend phases of kaolinite and carbonate. The evolutionary phase involves the chloritization phase of kaolinite in the presence of iron- and magnesium-supplied smectite illitization processes at a temperature between 165 and 200 °C [73], but reaction progress is inhibited due to the absence of kaolinite. At a lower temperature of about 120 °C and in the presence of carbonate minerals, another chloritization reaction via kaolinite and iron oxide could occur [12,74]. The possible explanation for hydrothermal chloritization cannot be discounted in the Mikambeni carbonaceous shale owing to the presence of Sagole thermal spring via the Tshipise fault [75,76]. This interpretation is consistent with the chloritization processes reported at the geothermal area of Iceland in which iron-rich smectite (saponite) is transformed into mixed-layer smectite-chlorite (S-C) at about 200 to 230 °C [50]. In support of the S-C mixed layer, corrensite is a resulting mineral composed of 50% of each chloride and smectite mineral [77]. With a further increase in temperature due to burial depth, chlorite crystallizes at temperatures greater than 240 °C due to a gradual replacement of iron-rich smectite with daughter chloride mineral in close zonation contact [78]. This reaction mechanism suggests a solid-state gradual replacement; however, the reaction pathway may be a continuous series transition instead of a stepwise progression [79] since shale has a low pore fluid to rock ratio due to poor interconnectivity of pores. Stepwise transition is noted to predominate in the system with high pore-fluid-to-rock ratio such as sandstone; hence, chloritization in the studied shale may be a by-product of smectite illitization. Although the rise in temperature of diagenetic chlorite could be promoted by contact diagenesis associated with the Sagole geothermal area, the pore fluid of Mikambeni shale is unaffected by the thermal spring due to extremely highly impermeable shale. Carbonate minerals commonly revert to the thermodynamically stable mineral as dolomite replaces limestones in the deep-burial diagenetic realm, as represented in Equation (7) [79]. Calcite dissolution increases with burial dolomitization in the presence of an organic-rich and interstitial fluid via stepwise precipitation of dolomite.

\[
\text{CaCO}_{3} + \text{Fe}^{2+} + \text{Mg}^{2+} \rightarrow 2\text{CaMg(CO}_{3})_{2} + 2\text{Ca}^{2+} \\
(7)
\]

The disappearance of calcite at relatively greater burial depths suggests a low pH medium, while its replacement by dolomite indicates an alkaline setting corresponding to oil window and organic matter maturation at a temperate range of 80–120 °C [80–82]. Promoted by sulfate reduction of pyrite mineral resulting from microbial activities, an increase in alkalinity medium favors precipitation of dolomite in the presence of Ca and Mg as burial depth increases [83]. The reduction process of \(\text{Fe}^{3+}\) to \(\text{Fe}^{2+}\) in the form of pyrite is evident from the framboidal structure revealed by SEM microscopy, thus suggesting a reducing or anoxic environment. Framboidal pyrites having bioclast structures that result from the reduction of organic matter in an organic-rich shale.

### 4.2. Methanogenesis Potentials of the Carbonaceous Shale

The biogenic methanogenesis from diagenetic carbonaceous shale may be attributed both to microbial degradation of organic matters and their thermal maturation as burial depth increases (Figure 7). The carboxyl C=O stretching absorption peak, around 1000–1220 cm\(^{-1}\), is stronger in spectra. The presence carboxyl group increases solubility properties. The post-depositional activities of microbial degradation on organic matter of shale deposit tend
to produce biogenic methane, sulfide gases, and organic acids fluid capable of dissolving some reactive mineral phases.

| Sedimentation and Mineral Changes | Mineral Diagenetic Effect | Organic Functional Group Progression | Structural Changes | Organic Matter Alteration | Pept. °C |
|----------------------------------|---------------------------|--------------------------------------|--------------------|---------------------------|----------|
| Fine-grained sediment, organic   | Bacterial mediation of     | Acetic                               | Grain-fabric,      | Biogenic and microbial    | 0        |
| plant material, inorganic minerals | organic matter,             | Protonic                             | ichno-rettic,      | gas \( \text{O}_2, \text{H}_2\text{S}, \text{CH}_4 \) | 10       |
| Terrigenous Carbonates           | interstitial brackish fluid |                                     | amorphous organic compound |                          | 20       |
| Kaolinitic alteration            | Oxallic                    | Dolomitic rhombs                     | Liquid H₂O         |                           | 50       |
| Feldspathic dissolution          |                           |                                       | Mixed liquid and gas |                           | 100      |
| Smectite                         |                           |                                       |                    |                           | 150      |
| Dolomitic replacement            | Phenolic                   | Frambooidal fibrous-lath like         | Thermal methane    |                           | 180      |
| Illitic crystallization          | Carboxylic                 | Lamellated                            | \( \text{CH}_4 \) |                           | 200      |
| Chloritic                        |                           |                                       |                    |                           | 2500     |

**Figure 7.** Methanogenesis potential of the studied carbonaceous shale.

Among the evidence of methanogenesis processes are the imprints of burrowing organisms, which suggest a connection with bioturbation processes capable of expelling biogenic gases from carbonaceous matter in shales [84]. The micro-FTIR revealed several linking bonds such as aromatic \( \text{C}=\text{C} \) stretching and amide group \( \text{N}=\text{H} \) bending in the acetic and propionic functional group, thus consistent with the organic macerals rich in protein, carbohydrate, lipids, and lignin [85]. At this stage, dewatering of pores first occurs at 10–50 °C, followed by biogenic methane from residual lignin at 50–70 °C (Figure 7) and characterized by carbonate minerals and acetic organic group. The water absorption spectra stretch between 3500 and 4000 cm\(^{-1}\) wavenumber band. This observation alludes to the carbon \( \delta^{13}\text{C} \) isotopic analysis, which reported an aceticlastic decomposition of aliphatic to aromatic components with the generation of early methane gas [86].

At greater depth, the organic functional group comprising mainly phenolic and carboxylic, which were observed alongside mineralogical components of illite and chlorite was associated with thermal methanogenesis. At depths greater than 2500 m, recent studies [87] have established the presence of diagenetic illite and chlorite minerals to characterized geochemical zones in which the temperature is above 180 °C and the low metamorphic zone. Although the Mikambeni shale is shallower, the proximity of Sagole and Siloam geothermal fields may play a role in the elevated temperature that caused the diagenetic and thermal maturation of organic matter. The alteration in I-M ordering from random to regular during montmorillonite illitization and chloritization in shales coincides with the temperature range of organic matter maturity and the onset of the oil and gas window in the Mikambeni shale. Depending on the nature of the organic matter, marine diatoms and fora-minifera undergo a catagenesis at the oil window, while the terrestrial higher plant materials exceed catagenesis to methanogenesis at about 200 °C, as indicated by the organic carboxylic group [87,88]. The methane is indicated by a strong absorption band around 3015 cm\(^{-1}\) and a band around 1305 cm\(^{-1}\).

The formation of solid carbon components involves a multi-stage process of kerogen degradation, initiated by deposition of organic matters [89]. Sources of organic matter could be either derived from the ocean, consisting of phytoplanktonic remains, or land, which consists mainly of vascular plant remains. However, accumulation of both marine and land-derived organic matter is common at deltas or floodplains of meandering rivers [90,91]. Under this oxic condition in the water column, the aerobic degradation of organic matter
continued until the dissolved oxygen was less than 0.2 mL/L H$_2$O [92]. The deficiency in oxygen may be due to climatic overprint, sea-level changes, tectonic activities, and hydrographic factors, resulting in an insufficient supply of oxygen to oxidize the organic materials [93,94]. In addition, the faster rate of organic matter accumulation in sediments more than the aerobic degradation and bacterial oxidation processes may contribute to the enrichment of organic carbon beyond its decomposition [95]. Below 0.2 mL/L H$_2$O oxygen concentration, anaerobic conditions and degradation of organic matter begin, giving rise to the solid carbon-containing components in shale sediments [95]. This anoxic setting is established as all the oxidants in the shale sediments deplete due to further reduction processes caused by increasing burial depth, promoting the activities of sulfide-reducing bacteria [96]. Studies [97–100] have highlighted minerals and compounds such as pyrite, sphalerite, hydrogen sulphide, and iron sulfide as evidence of the reduction process that ensued from sulfide-driven bacteria. With further increase in burial depth below the sediment–water interface, the methanogens begin to degrade the organic carbon to generate biogenic methane. This is consistent with the previous work of Tourtelot (1979), which argued that methane is unlikely to form unless all sulphates are depleted while vanadium and nickel are concentrated.

5. Conclusions

The combination of FTIR, SEM-EDX, and mineralogical analyses demonstrated a promising approach to assess methanogenesis potentials in carbonaceous shale rock. In shale rock that is dominated by terrigenous higher plants, the biogenic methanogenesis is found to correspond to the absorption spectrum stretching from 1650–1220 cm$^{-1}$ frequencies. This spectrum reflects the C-O stretching and OH deformation of acetic and phenolic groups in all studied samples. Given the strong absorptions around 2000–2300 cm$^{-1}$, CO$_2$ is produced by decarboxylation of organic matter. The gas becomes acidified and dissolves feldspar to release K$^+$ ions, which promoted the illitization of smectite. The SEM microscopy depicted a framboidal structure, which indicated a sulfate reduction of pyrite mineral. The reduction process results from microbial activities in an anoxic milieu and causes an increase in alkalinity medium that favors precipitation of dolomite in the presence of Ca and Mg as burial depth increases. The contact diagenesis from the proximity of Sagole geothermal spring via Tshipise fault is suggested to have enhanced the transformation of smectite to chlorite via a mixed-layer corrensite in a solid-state gradual replacement reaction pathway. The presence of diageneric chloride mineral is characteristic of low-grade metamorphism or high diagenetic zone at a temperature around 200–230 °C and corresponds to thermal breakdown of kerogen to methane at a strong absorption band around 2850–3000 cm$^{-1}$.

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