Diagenetic influences on iron-bearing minerals in Devonian carbonate and siliciclastic rocks of Estonia

Alla Shogenova and Anne Kleesment

Institute of Geology at Tallinn University of Technology, Ehitajate tee 5, 19086 Tallinn, Estonia; alla@gi.ee

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Abstract. The Estonian Devonian sediments consist of carbonate, siliciclastic, and mixed carbonate-siliciclastic deposits that have undergone extensive dolomitization. Chemical analysis of 165 samples and XRD mineralogical analysis of 10 samples from 8 drill cores were carried out and combined with previous results, obtained particularly on the heavy and clay fraction residues of 131 samples. The carbonate residues include illite, illite-smectite, and chlorite, with accessory biotite, muscovite, pyrite, goethite, hematite, siderite, sphalerite, and magnetite. The residues of siliciclastic rocks consist of quartz, K-feldspar, illite, montmorillonite-chlorite and chlorite, with accessory biotite, muscovite, leucoxene, ilmenite, pyrite, hematite, goethite, and siderite. The iron content of rocks is related to the detrital input during primary sedimentation or diagenetic products formed during cementation, dolomitization, and authigenic mineral growth. Iron minerals underwent chemical alteration during diagenesis and are partly corroded and dissolved. Fe(III) minerals, responsible for red coloration of rocks, became dominant due to oxidation, low water table, and arid climate, which prevailed during diagenesis. The red coloration of siliciclastic rocks is due to hematite coatings of quartz grains. It may be primary in origin, but also diagenetic due to alterations of maghemite to hematite and dispersed distribution of Fe-oxides in dolomite cement. The red coloration of carbonate rocks (dolostones and marlstones) may have an early diagenetic origin prior to dolomitization, with possible redistribution of colour at later diagenetic stages. The variable origin of iron minerals (primary, early, middle, and late diagenetic) is distinguishable by thin-section petrography, and can be also supported by chemical and mineralogical data. The average \( \frac{Fe_2O_3_{\text{total}}}{Al_2O_3} \) ratio in the studied carbonates is higher than in siliciclastics, being in accordance with the distribution of these elements in the Earth’s crust. In contrast to the Earth’s crust data, the average TiO2/Al2O3 and K2O/Al2O3 ratios are higher in Devonian siliciclastics. This could be explained by alteration of clay during late diagenesis.

Key words: Devonian, diagenesis, carbonate rocks, siliciclastic rocks, mixed carbonate-siliciclastic rocks, iron minerals, insoluble residue, Baltic basin.
INTRODUCTION

This is the first of the two papers dealing with the chemico-mineralogical influence of diagenetic processes on Devonian carbonate and siliciclastic rocks of Estonia. Here the results of previous geochemical, mineralogical, and petrographical studies are supplemented by new data from eight drill cores (Fig. 1). A second forthcoming article will consider the rock magnetic properties of the same sequences. The initial study (Shogenova 1999) revealed the importance of diagenetic processes in Estonian Palaeozoic rocks, but the problems are wide-ranging in both space and time (Haese et al. 1998; Haubold 1999; Marton 1999; Wilson & Roberts 1999). Recent interest has focused on processes related to diagenesis (Passier et al. 2001; Lewchuk et al. 2002; Schneider et al. 2004a, b). The main event of the Hercynian orogeny in Europe was the consolidation of mountain-fold belts. As the Baltic Devonian sedimentary basin was located in the northwestern part of the East European Craton, it lay in the equatorial region of the Laurussia continent that formed at the end of the Silurian (Ziegler 1988). Both tectonic movements and eustatic sea-level oscillations influenced the development of sedimentary sequences in this region (Ziegler 1988; Mokrik 2003). Epicontinental shallow sea sediments have a complex cyclic structure in which siliciclastic and carbonate sediments alternate recurrently. Pure dolostone, dolomitic marlstone, siliciclastic sediments, and mixed carbonate-siliciclastic rocks are represented in southern and southeastern Estonia (Kleesment & Shogenova 2005). The thickness of the Devonian sequence in the eight studied boreholes (Fig. 1) is from 111 m in the Häädemeeste core up to 352 m in the Võru core (Kleesment & Shogenova 2005).

Fig. 1. Distribution of Devonian rocks (grey shading) and location of the sampled boreholes.
The main influx of siliciclastic material was from the Scandinavian granite
massifs (Kuršs 1992; Kleesment 1997; Plink-Björklund & Björklund 1999), while
the carbonate sediments are mostly represented by dolostones. The Devonian
sediments overlie with a stratigraphical gap the Silurian or Ordovician (Kleesment
& Shogenova 2005). The Estonian part of the Baltic Devonian basin is unique
as here the alternating carbonate, siliciclastic, and mixed carbonate-siliciclastic
rocks have been influenced by subsequent diagenetic dolomitization episodes
during which a number of minerals (feldspar, pyrite, goethite, gypsum) were
formed. These and primary minerals were later partly corroded, dissolved, and
recrystallized by diagenetic fluids (Kleesment 1984, 1998). The sediment sequences
of the area thus contain mixed rocks of primary and secondary genesis (Kleesment
& Shogenova 2005). Some of iron-bearing minerals, of primary and diagenetic
origin, are responsible for colours ranging from pink to red and red-grey mottled
varieties that also vary between carbonate and siliciclastic rocks (Fig. 2). Several
authors have studied diagenesis in the southern part of the Baltic Devonian basin
(e.g. Narbutas 1981; Šimškaite & Simkevičius 1981). The influence of different
iron-bearing minerals has been interpreted considering the red colouring of
Devonian siliciclastic rocks (Kuršs & Stinkule 1972). Diagenetic replacement
of magnetite by goethite in siliciclastic rocks has mostly been described in sections
with thin intercalations of siliciclastic and clayey-dolomitic rocks (Paluskova
1988; Abdalla et al. 1997). The replacement of ilmenite by leucoxene during
diagenesis has been known for a long time, but is often ascribed to different
stages from early diagenesis in mildly reducing and neutral-acid tropical conditions
(Weibel 2003) to late diagenesis (Al-Juboury et al. 1994). The alteration of
ilmenite is an important source of iron for diagenetic minerals (Pe-Piper et al.
2005). As published data on the chemical composition of the Baltic Devonian
rocks are scarce, here mainly the results of Shogenova et al. (2003a, b) and
Kleesment & Shogenova (2005) are used. The influence of dolomitization on the
composition and properties of Estonian carbonates has earlier been studied in
However, this phenomenon also occurs in the Devonian of southern Estonia
and possibly all these areas are affected by Devonian dolomitization fluids. An
increase in the total iron content and magnetic parameters has been recorded in
the Ordovician, and to some extent Silurian dolostones in comparison with lime-
stones of these ages. The influence of dolomite cement on siliciclastic rocks has
been described for Cambrian rocks of Estonia (Shogenova et al. 2001) and in the

Diagenesis is defined as a series of chemical, physical, and biological processes
leading to a significant change in sediments, which started after their deposition
and continued during and after their lithification (Larsen & Chilingar 1979;
Morse & Mackenzie 1990). Carbonate diagenesis includes such major processes
as cementation, microbial micritization, neomorphism, dissolution, compaction,
and dolomitization and it operates in three main environments: marine, near-
surface meteoritic, and burial (Tucker & Wright 1994). Diagenesis of siliciclastic
Lithology, colour, and gamma-ray log of the Mehikoorma (421) drill core. The total iron content (filled circles) of core samples and their Fe₂O₃/FeO ratio (open triangles) are shown in a log-decimal scale. In addition, the Fe₂O₃total/Al₂O₃ ratio is shown for siliciclastic rocks (open circles), mixed rocks (filled diamonds), and carbonate rocks (crosses). The Devonian is represented by the top of the Emsian Stage (Rezekne Regional Stage, D₁rz), Eifelian Stage including the Pärnu (D₂pr), Narva (D₂nr), and Aruküla (D₂ar) regional stages, and the oldest part of the Givetian Stage – Burtnieki Regional Stage (D₂br). The Narva Regional Stage is subdivided into the Vadja (D₂nrV), Leivu (D₂nrL), and Kernave (D₂nrK) formations and the Aruküla Regional Stage into the Viljandi (D₂arvl), Kureküla (D₂arkr), and Tarvastu (D₂artr) members (Kleesment & Shogenova 2005).

rocks includes compaction, cementation, reworking, authigenesis, replacement, recrystallization, leaching, hydration, bacterial actions, and formation of concretions.

The division of diagenesis most widely accepted in the scientific literature has three stages: early (immediately after deposition), middle (deep burial), and late (during and after uplift) (Fairbridge 1967). Late diagenesis was followed by weathering (Larsen & Chilingar 1979).
In case of siliciclastic rocks, because of oxidation and reduction reactions, diagenesis is subdivided into the redoxomorphic, locomorphic, and phyllomorphic stages. The redoxomorphic stage is mostly responsible for the bulk final colour of the rock, while later changes in rock colour are smaller. During the second stage primary cementation develops and the rock becomes lithified. Authigenic overgrowth and clay alteration take place during the third stage (Dapples 1979).

During the redoxomorphic stage iron is moved into the ferrous state and sulphur is fixed as pyrite, which is scattered throughout the rock. If an organic fraction is significant, the rock has grey colour (Love 1971). Red colour may appear in oxidizing conditions when detrital iron oxides or clay minerals are transformed into hematite and related ferric oxides and hydrates, occurring in the form of films, matrix or pore filling.

DATA AND METHODS

The bulk chemical composition (SiO$_2$, Al$_2$O$_3$, TiO$_2$, K$_2$O, Na$_2$O, Fe$_2$O$_3$total, CaO, MgO, MnO, and P$_2$O$_5$ contents) of 165 samples from eight boreholes was determined by X-ray fluorescence analysis in the All-Russian Geological Institute, St. Petersburg, using the method described in Teedumäe et al. (2006). The insoluble residue (IR) and FeO contents were measured by wet chemical analysis at the Institute of Geology at Tallinn University of Technology. The Pearson correlation was calculated for the pairs of measured parameters, without logarithmic transformation.

Chemical analyses

The IR was determined gravimetrically using carbonate wet chemical analysis. An amount of 50 mL of HCl (1 : 4) was added to 0.5 g of rock powder and dried in a water bath. The remaining residue was mixed with 5 mL of concentrated HCl and 30–40 mL of hot water (90–100°C), filtrated and washed with hot water (90–100°C). The filtrated residue was heated to 1000°C for one hour and then weighed. The concentration of ferrous oxide (FeO) was determined photocolorimetrically with α-α dipirisol (C$_{12}$H$_8$N$_2$). Fe$^{2+}$ forms with α-α dipirisol an orange-red water soluble complex with pH = 4. It is stable in respect to oxygen and over long time periods. The colour appears very fast and depends strongly on Beer’s law for wide ranges of Fe$^{2+}$ concentrations. The maximum photoabsorbability occurs at 505 nm wavelength.

Silicate rocks and minerals were decomposed by a mixture of HF and H$_2$SO$_4$. The solution was boiled for 5 min so that the carbonate rocks were decomposed by H$_2$SO$_4$. The value pH = 4 was controlled by the H$_3$BO$_3$ solution. This method allows the FeO concentration in the range of 0.1–20% to be determined in samples with a mass between 0.1 and 0.5 g.
Mineralogical analysis

Mineralogical analysis of the clay fraction and heavy minerals has earlier been carried out in the same boreholes, though on different samples. The mineralogical composition of the clay fraction of 81 samples from three boreholes was studied using the URS-50 IM and Dron-1 diffractometers, allowing the identification of the main minerals with an accuracy of about 5% (Fe-filtered Co- and Cu-radiation; Utsal 1971; Stinkule & Utsal 1975). The mineralogical composition of heavy minerals in 131 samples from four boreholes was determined in immersion liquids under a microscope using plane-polarized, transmitted light. Samples with dolomite cement were crushed and dissolved in 10% acetic acid and IR grains with a diameter of <0.01 mm were removed by washing. Grains >0.01 mm were separated by sieving and studied in transmitted and reflected light under the binocular (BM) and petrographic microscope (PM). The 0.05–0.1 mm fraction was separated using bromoform and the light and heavy minerals were determined separately by the immersion method under the PM in plane and cross-polarized light. About 300–500 grains were counted for each mineral spectrum and expressed in per cent (Kleesment & Valiukevičius 1998; Kleesment 2001, 2003). Ilmenite, magnetite, pyrite, and garnet were determined by microscopical analysis. Grains >0.1 mm were investigated using the BM. The content of magnetite was estimated roughly with a magnet and, in some samples, the occurrence of magnetite was checked using XRD techniques. Pyrite was identified in thin sections and in immersion liquid. In addition, during the present study the bulk mineral composition of 10 samples was determined using the HZG4 diffractometer (Fe-filtered Co-radiation). For these analyses sample powders were mixed with alcohol and spread on glass slides. A range from 5 to 40° 2θ was step-scanned (step 0.05° 2θ, counting rate 3 s). Additionally 55 thin sections were studied under the PM with up to 200 times magnification.

RESULTS

Composition of the studied rocks

Chemical composition

The samples can be subdivided into four lithological groups based on chemical parameters (Kleesment & Shogenova 2005). As the carbonate rocks in the eight boreholes considered are almost fully dolomitized, four general rock types could be identified using only one chemical parameter – IR. From 165 rock samples, the following rock types were obtained (Fig. 3, Table 1):
(1) dolostones (IR < 25%) – 40 samples;
(2) dolomitic marlstones (25% < IR < 50%) – 29 samples;
(3) mixed carbonate-siliciclastic rocks (50% < IR < 70%) – 39 samples represented by dolostones, dolomitic marlstones, siltstones, and sandstones;
(4) siliciclastic rocks (IR > 70%) – 57 samples represented by cemented sandstones, siltstones, and mudstones.
Fig. 3. MgO content by XRF data versus IR content by wet chemical analysis. Mixed and siliciclastic rock samples, located away from the correlation line, include gypsum and calcite cement.

The first type includes dolostones from pure dolomite to strongly argillaceous rock, with an IR content between 2.7 and 23.5% and MgO content of 14.5–19.9% (Table 1). Dolostones had the lowest average total iron content (1.56%), but the highest ratios of Fe$_2$O$_3$/Al$_2$O$_3$, Fe$_2$O$_3$/K$_2$O, and Fe$_2$O$_3$/TiO$_2$ (Table 1, Figs 2, 4a–c). The content of oxides (Al$_2$O$_3$, K$_2$O, TiO$_2$) in the IR, which reflects the clay content of the rocks, SiO$_2$ content, total iron content, and K$_2$O/Al$_2$O$_3$ and TiO$_2$/Al$_2$O$_3$ ratios were the lowest in dolostones in comparison with the other rock types. The average FeO content (0.16%) and Fe$_2$O$_3$/FeO ratio were also the lowest among the studied rock types. The FeO content of dolostones was not dependent on the total iron and TiO$_2$ contents (Table 1, Fig. 4d–f).

Dolomitic marlstones contain 25.9–49% IR and 10.3–16% MgO. Total iron, Al$_2$O$_3$, K$_2$O, and TiO$_2$ contents, and K$_2$O/Al$_2$O$_3$ and TiO$_2$/Al$_2$O$_3$ ratios were higher than in dolostones, while Fe$_2$O$_3$/Al$_2$O$_3$, Fe$_2$O$_3$/K$_2$O, and Fe$_2$O$_3$/TiO$_2$ ratios were lower (Table 1, Fig. 4a–c). The FeO content was on average (0.23%) higher than in dolostones, but the Fe$_2$O$_3$/FeO ratio was in the same limits (Table 1; Figs 2, 4d–e). The Fe$_2$O$_3$/Al$_2$O$_3$, Fe$_2$O$_3$/K$_2$O, and Fe$_2$O$_3$/TiO$_2$ ratios in carbonate rocks (dolostones and dolomitic marlstones) were higher than in mixed and siliciclastic rocks (Table 1; Figs 2, 4a–c). The correlation of the total iron content with Al$_2$O$_3$, K$_2$O, and TiO$_2$, contents of carbonate rocks (dolostones and dolomitic marlstones) taken together was relatively high (correlation coefficients 0.84, 0.76, and 0.82, respectively), but the correlation was negative with MgO and CaO contents (−0.63 and −0.7; Fig. 4d).

Siliciclastic rocks are represented by sand- and siltstones with cement consisting of clay, dolomite and iron-bearing minerals. They contain 70.2–97.6% IR and 0.14–6.6% MgO. Three siliciclastic samples which lie away from the correlation line (Fig. 3) contained calcite (Värskka (6) borehole) and gypsum cement (Värskka (6)
Table 1. Chemical composition of rocks*

<table>
<thead>
<tr>
<th>Studied parameter</th>
<th>Dolostones</th>
<th>Dolomitic marlstones</th>
<th>Mixed carbonate-siliciclastic rocks</th>
<th>Siliciclastic rocks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Insoluble residue, %</td>
<td>2.7–23.5/13.2</td>
<td>25.9–49/34.3</td>
<td>51–69.4/59.5</td>
<td>70.2–97.6/83</td>
</tr>
<tr>
<td>MgO, %</td>
<td>14.5–19.9/17.7</td>
<td>10.3–16/13.3</td>
<td>6–11/8.4</td>
<td>0.14–6.6/3.1</td>
</tr>
<tr>
<td>CaO, %</td>
<td>22–31.7/26.3</td>
<td>13.2–19.9/18.2</td>
<td>11–16.1/10.5</td>
<td>0.15–10.8/3.9</td>
</tr>
<tr>
<td>SiO₂, %</td>
<td>0.74–17.5/10</td>
<td>19.1–44.3/27.6</td>
<td>36.1–66.5/49.8</td>
<td>53.6–94.7/71.3</td>
</tr>
<tr>
<td>Al₂O₃, %</td>
<td>0.55–5.7/2.9</td>
<td>1.94–12/6.24</td>
<td>1.11–16.8/7.5</td>
<td>1.64–16.6/7.7</td>
</tr>
<tr>
<td>K₂O, %</td>
<td>0.2–1.9/1</td>
<td>1.4–3.9/2.5</td>
<td>0.9–6.5/3.6</td>
<td>1.22–8.7/4.5</td>
</tr>
<tr>
<td>TiO₂, %</td>
<td>&lt;0.005–0.26/0.11</td>
<td>0.14–0.47/0.31</td>
<td>0.01–0.8/0.4</td>
<td>0.06–0.99/0.46</td>
</tr>
<tr>
<td>K₂O/Al₂O₃</td>
<td>0.08–0.48/0.36</td>
<td>0.24–0.7/0.43</td>
<td>0.3–0.87/0.55</td>
<td>0.38–0.78/0.62</td>
</tr>
<tr>
<td>TiO₂/Al₂O₃</td>
<td>0.001–0.05/0.04</td>
<td>0.035–0.08/0.05</td>
<td>0.003–0.09/0.05</td>
<td>0.022–0.18/0.06</td>
</tr>
<tr>
<td>Fe₂O₃total, %</td>
<td>0.63–3.3/1.56</td>
<td>0.9–5.5/2.62</td>
<td>0.4–7.5/3.1</td>
<td>0.45–9.1/2.57</td>
</tr>
<tr>
<td>TiO₂/Al₂O₃</td>
<td>0.011/0.012</td>
<td>0.0011/0.012</td>
<td>0.0001/0.001</td>
<td>0.011/0.012</td>
</tr>
<tr>
<td>FeO, %</td>
<td>0.03–0.43/0.16</td>
<td>0.03–0.43/0.23</td>
<td>0.01–0.94/0.24</td>
<td>0.05–0.52/0.19</td>
</tr>
<tr>
<td>Fe₂O₃/FeO</td>
<td>2.5–51/13.7</td>
<td>2.6–50/14</td>
<td>3.6–93/19</td>
<td>1–41/15.5</td>
</tr>
<tr>
<td>Fe₂O₃total/Al₂O₃</td>
<td>0.29–3.15/0.68</td>
<td>0.21–0.91/0.43</td>
<td>0.11–0.71/0.41</td>
<td>0.1–0.58/0.31</td>
</tr>
</tbody>
</table>

* Five samples with calcite and gypsum cement are not included in the table. Std. Dev., standard deviation.

and Võru boreholes). The total iron content varied largely (0.45–9.1%), however, the average value (2.6%) was close to that of marlstones and mixed rocks (Fig. 4a–d, Table 1). The Al₂O₃, K₂O, and TiO₂ contents of the siliciclastic rocks and the K₂O/Al₂O₃ and TiO₂/Al₂O₃ ratios were the highest among the studied rock types (Fig. 4a–c, Table 1). Correlation of the total iron content with Al₂O₃, K₂O, and TiO₂ was also high (coefficients 0.89, 0.68, and 0.88, respectively), but there was no significant correlation with carbonate components (CaO and MgO; Fig. 4a–d). Aluminium oxide has entered mainly clay minerals, titanium oxide reflects titanium minerals, and potassium oxide has entered both clay minerals and K-feldspar in these rocks.
The Fe$_2$O$_3$$_{\text{total}}$/Al$_2$O$_3$, Fe$_2$O$_3$$_{\text{total}}$/K$_2$O, and Fe$_2$O$_3$$_{\text{total}}$/TiO$_2$ ratios in siliciclastic rocks were lower than in carbonate and mixed rocks. The FeO content was 0.05–0.52% and had significant, but not high correlation ($R = 0.37$) with the total iron content. The Fe$_2$O$_3$$_{\text{total}}$/FeO ratio had the narrowest limits, but on average it was higher than in carbonate rocks.

**Mixed rocks** include 51–69.4% IR and 6–11% MgO. The limits of the total iron content in this group (0.4–7.5%) were wider than in marlstones and narrower than in siliciclastic rocks, while the average (3.1%) was higher than in the other rock groups (Table 1). The Fe$_2$O$_3$$_{\text{total}}$/Al$_2$O$_3$, Fe$_2$O$_3$$_{\text{total}}$/K$_2$O, and Fe$_2$O$_3$$_{\text{total}}$/TiO$_2$ ratios were lower than in carbonate, but higher than in siliciclastic rocks (Fig. 4a–c). The Fe$_2$O$_3$/FeO ratio was the highest among the studied rock types. The TiO$_2$, and Al$_2$O$_3$ contents varied widely (Table 1; Figs 2, 4a,c), while their means and the TiO$_2$/Al$_2$O$_3$ ratio were close to those of dolomitic marlstones. The K$_2$O content (0.9–6.5%; Fig. 4b) and average K$_2$O/Al$_2$O$_3$ ratio were higher than in dolomitic marlstones, but lower than in siliciclastic rocks (Table 1). Correlation of the total iron content with Al$_2$O$_3$, K$_2$O, and TiO$_2$ contents was the highest among all rock groups (coefficients 0.93, 0.8, and 0.87, respectively; Fig. 4a–c). The correlation coefficients of FeO with total iron ($R = 0.68$) and TiO$_2$ contents ($R = 0.61$) were higher than for siliciclastic and carbonate rocks.

**Lithology and petrography**

Dolostones are light grey to light brownish-grey. The matrix of dolostone is cloudy, and dominantly aphanocrystalline (<0.004 mm), with rare very fine crystalline grains (0.005–0.01 mm). The structure is often patchy, due to different clay components and pigmentation from iron minerals. Scattered crystals of pyrite and goethite/hematite (both 0.001–0.005 mm) were found in the matrix (Fig. 5a,b). The IR was mainly represented by a muddy fraction (<0.01 mm) consisting predominently of illite accompanied by a considerable amount of chlorite. Mixed layers of montmorillonite-chlorite, illite-smectite, illite-chlorite, and illite-montmorillonite also occurred. Dolomitic marlstones are grey, often with violet or greenish shades, mottled reddish-brown (Fig. 5c,d) to greenish-grey, mostly very fine to fine-grained (0.004–0.01 mm), cloudy or semitransparent (Fig. 5c,d) and alternating with clay- and dolostone. The structure of the beds is commonly massive. The matrix is pigmented with indistinct patches of hematite or goethite, containing also fine and varied detrital grains. The IR was mainly represented by muddy particles (<0.01 mm). The muddy fraction was characterized by a high content of illite (60–80%) accompanied with chlorite. Sometimes an admixture of mixed-layer chlorite-montmorillonite was found (up to 20%). The content of detrital particles (>0.01 mm) was 5–20%. The general mineralogical composition of reddish-brown, grey, and pink loosely and carbonate cemented siliciclastic rocks is similar. Quartz accompanied by K-feldspars and mica minerals dominates (Kleesment & Shogenova 2005). A high variability was observed in the content of accessory minerals. During diagenesis of loosely cemented siliciclastic rocks
Fig. 4. (a) Total iron content on a log-decimal scale versus $\text{Al}_2\text{O}_3$ content. The correlation coefficient $R$ of the total iron content and $\text{Al}_2\text{O}_3$ content is 0.84 for dolostones and dolomitic marlstones, $R = 0.93$ for mixed rocks, and $R = 0.89$ for siliciclastic rocks. (b) Total iron content on a log-decimal scale versus $\text{K}_2\text{O}$ content. $R = 0.76$ for dolostones and dolomitic marlstones, $R = 0.80$ for mixed rocks, and $R = 0.68$ for siliciclastic rocks. (c) Total iron content on a log-decimal scale versus $\text{TiO}_2$ content. $R = 0.82$ for dolostones and dolomitic marlstones, $R = 0.87$ for mixed rocks, and $R = 0.88$ for siliciclastic rocks. (d) Total iron content versus $\text{MgO}$ content. $R = -0.63$ for carbonate rocks, and is insignificant for dolomitic marlstones, mixed rocks, and siliciclastic rocks. (e) FeO content on a log-decimal scale versus total iron content. $R = 0.68$ for mixed rocks, $R = 0.37$ for siliciclastic rocks, and is insignificant for dolostones and dolomitic marlstones. (f) FeO content versus $\text{TiO}_2$ content. $R = 0.35$ for siliciclastic rocks and $R = 0.61$ for mixed rocks, and is insignificant for dolostones and dolomitic marlstones.
detrital magnetite was replaced by goethite (Kleesment 1984). Cemented siliciclastic rocks have mostly a dolomite matrix. Only in a few cases calcite cement occurred in the lower part of the sequence in East Estonia (Võru core). Some gypsum cemented layers were observed in the Vadja Formation in the Värska (6) core. Usually cement formed 30–40% (Fig. 6a,b), more rarely 10–30% of the rock.
Patchy distribution of cement was common, with grain-supported and cement-supported spots. Pigmentation of the matrix by goethite or hematite was either patchy or appeared along bedding planes (Fig. 6a,b). Occasionally pyrite cement was found.

Varicoloured mixed carbonate-siliciclastic rocks are represented by cement-supported silt- and sandstone, and by dolomitic marlstone and dolostone including siliciclastic material (Fig. 6c). Two mixed samples which do not lie on the correlation line (Fig. 3) had the MgO and CaO contents of 2.7 and 17.9% and 6.3 and 16.6%, respectively. They represent sandstones with calcite and patchy pyrite cement. While mixed carbonate and siltstone samples were formed during sedimentation, mixed dolomitic sandstone, in some cases also dolomitic coarse-grained siltstone, are in this group based on diagenetic influences. The carbonate content of mixed rocks was 30–50% (Fig. 6d). Cement consists of cloudy fine-crystalline or medium- to coarse-crystalline clear dolomite. Rare fine interlayers with calcite cement were observed. Patchy distribution of cement was common, which is often associated with pyrite. Detrital grains and crystals of pyrite were often coated with carbonate rims (Fig. 6d). This indicates that pyrite has formed earlier than carbonate. In rare cases calcite occurred as clear medium- to coarse-crystalline grains.

Mineralogical composition

Bulk mineralogical qualitative XRD analysis showed that dolostones were comprised mainly of dolomite and IR including quartz and K-feldspar (Fig. 7a) and very small amounts of illite, illite-smectite, chlorite, and kaolinite. Dolomitic marlstones are composed of the same minerals as dolostones. In some samples hematite was also recorded. Mixed carbonate-siliciclastic rocks consist of quartz, dolomite, illite, chlorite, and K-feldspar (Fig. 7b). Siliciclastic rocks consist of quartz, K-feldspar, dolomite, illite, and chlorite (Fig. 7c). Mineralogical XRD analysis of a bulk sample can only determine the main components in the rock; occasionally an admixture of iron-bearing minerals is also recorded. The XRD mineralogical analysis of the clay fraction (<0.005 mm) is more efficient for identifying the iron-bearing minerals (Table 2, Fig. 8a), since most of them occur in the clay fraction of the rocks. The clay fraction of dolostones and dolomitic marlstones which formed 0.8–60.1% of the rock in 22 samples, mainly consists of illite, chlorite, K-feldspar, and quartz, with an admixture of goethite, hematite, and siderite. The clay fraction of siliciclastic rocks, forming 0.3–69% of the rock in 59 samples, mainly consists of illite, chlorite, kaolinite, montmorillonite-chlorite, feldspar, quartz and smaller amounts of hematite, goethite, and siderite (Table 2). The clay fraction content was higher in carbonate rocks than in siliciclastic rocks, but average illite, chlorite, montmorillonite, and siderite contents were similar. The kaolinite, goethite, and hematite contents were higher in siliciclastic rocks, while the quartz and feldspar contents were higher in the clay fraction of carbonate rocks (Fig. 8a).
Fig. 5. Thin sections of dolostones and dolomitic marlstones. (a) Aphanocrystalline cloudy dolostone penetrated by vugs and fractures filled with authigenic transparent dolomite (D). Abundant fine goethite (hematite) grains and pyrite crystals (5–15 µm) are scattered in the matrix. Concentrations of goethite (hematite) grains occur in places (F). Mehikoorma (421) core, 190.6 m, plane-polarized light. MgO content 18.5%, IR content 8.9%, Fe₂O₃total content 1.63%, Fe₂O₃total/Al₂O₃ ratio 1.78. For mineralogical analysis see Fig. 7a. (b) Aphanocrystalline semitransparent dolostone (<0.004 mm). Abundant Fe-hydroxide (F) grains (0.005–0.01 mm) and pigmented spots with irregular boundaries (0.01–0.02 mm) dispersed in the dolomite matrix. The rock is penetrated by wavy lense-shaped planes connected with vugs and in some places by aggregates of authigenic concentric crystals of sphalerite (S) (0.1–0.2 mm). Häädemeeste core, 79.0 m, Narva Stage, Vadja Formation. Plane-polarized light. MgO content 19.5%, IR content 3.34%, total iron content 1.73%, Fe₂O₃total/Al₂O₃ ratio 3.15. (c) Very finely to finely crystalline silty dolomitic marlstone, patchily pigmented by Fe-oxides (F). Pigmentation is strong in places, and irregular opaque patches of hematite/goethite occur. Mehikoorma (421) core, 161.2 m, plane-polarized light. MgO content 18.5%, IR content 10.8%, total iron content 3.91%, Fe₂O₃total/Al₂O₃ ratio 0.54. (d) Cloudy very finely crystalline dolomitic marl, patchily pigmented by Fe-hydroxides (F). Taagepera core, 175 m, plane-polarized light. MgO content 11.3%, IR content 41.2%, total iron content 5.52%, Fe₂O₃total/Al₂O₃ ratio 0.46.
Fig. 6. Thin sections of siliciclastic and mixed carbonate-siliciclastic rocks. (a) Very fine-grained wavy-bedded sandstone. Argillaceous dolomite is pigmented by Fe-oxides (F). Strong pigmentation occurs along bedding surfaces. Valga (10) core, 118.5 m, plane-polarized light. IR content is 93%, of which clay minerals form about 10%. Among detrital part quartz dominates (80%), feldspars and micas form both about 10%. Total iron content 2.94%, $\text{Fe}_2\text{O}_3/\text{Al}_2\text{O}_3$ ratio 0.32. (b) Siltstone, indistinctly wavy-bedded, with patchy goethite/hematite pigmentation (F). Tartu (453) core, 46.0 m, crossed polars. IR content is 72.4%, of which clay minerals form about 40%. Among detrital grains quartz forms about 70%, feldspars form 20–25%, and micas 5–10%. Total iron content 6.47%, $\text{Fe}_2\text{O}_3/\text{Al}_2\text{O}_3$ ratio 0.49. (c) Mixed dolomite-siliciclastic rock: wavy-bedded silty dolomitic marlstone. Strong hematite-pigmentation (F) alternating with less pigmented beds (0.2–0.9 mm). Mehikoorma (421) core, 154.6 m, plane-polarized light. IR content is 53.2%, of which clay minerals constitute about 35%. Among detrital minerals quartz make up approximately 70%, feldspars and micas account both for 15%. Total iron content 3.21%, $\text{Fe}_2\text{O}_3/\text{Al}_2\text{O}_3$ ratio 0.39. (d) Fine- to medium-grained cement-supported sandstone with clear calcite cement. Pyrite (P), whose crystals are surrounded by calcite rims. Värskas (6) core, 293.0 m, crossed polars. IR content is 64.1%, of which clay minerals constitute about 5%. Among detrital minerals quartz dominates, making up 85–90% of the rock. Feldspars make up the remaining 10–15%. Total iron content 2.3%, $\text{Fe}_2\text{O}_3/\text{Al}_2\text{O}_3$ ratio 1.03.
Fig. 7. Bulk mineralogical composition of (a) dolostone of the Mehikoorma (421) core, 190.6 m, (b) silty dolomitic marlstone (mixed rock) of the Häädemeeste core, 33.8 m, and (c) sandstone of the Vär ska (6) core, 125.5 m. XRD analysis by T. Kallaste. Q, quartz; K-f., K-feldspar; Kl, kaolinite; I, illite; Dol., dolomite; Hem., hematite.
Table 2. Mineralogical composition of the clay fraction (<0.005 mm) of the rocks of the Mehikoorma (421), Ruhnu (500), and Võru cores, determined by XRD analysis

<table>
<thead>
<tr>
<th>Formation, rocks</th>
<th>Number of samples</th>
<th>Statistical parameter</th>
<th>Dolomitic component of rock</th>
<th>Weight per cent of clay fraction</th>
<th>Clay component, %</th>
<th>Other minerals, %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Illite</td>
<td>Chlorite</td>
</tr>
<tr>
<td>Aruküla, siliciclastic</td>
<td>19 Min</td>
<td>0</td>
<td>1.4</td>
<td>52</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Kernave, siliciclastic</td>
<td>Max</td>
<td>30.8</td>
<td>69</td>
<td>93</td>
<td>46</td>
<td>25</td>
</tr>
<tr>
<td>Avg</td>
<td>6.3</td>
<td>21.3</td>
<td>81.5</td>
<td>14.3</td>
<td>3.8</td>
<td>0.4</td>
</tr>
<tr>
<td>Leivu, siliciclastic</td>
<td>9 Min</td>
<td>0</td>
<td>1</td>
<td>55</td>
<td>18</td>
<td>0</td>
</tr>
<tr>
<td>Avg</td>
<td>37.1</td>
<td>54.1</td>
<td>82</td>
<td>40</td>
<td>7</td>
<td>0.8</td>
</tr>
<tr>
<td>Leivu, carbonate</td>
<td>18 Min</td>
<td>3</td>
<td>0.8</td>
<td>48</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>Vadjá, carbonate</td>
<td>Max</td>
<td>72.9</td>
<td>60.1</td>
<td>88</td>
<td>42</td>
<td>0</td>
</tr>
<tr>
<td>Avg</td>
<td>36</td>
<td>36</td>
<td>69.7</td>
<td>22.9</td>
<td>0</td>
<td>1.6</td>
</tr>
<tr>
<td>Pärnu, siliciclastic</td>
<td>31 Min</td>
<td>0</td>
<td>0.3</td>
<td>15</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Rezekne, siliciclastic</td>
<td>Max</td>
<td>38.2</td>
<td>37</td>
<td>99.8</td>
<td>37</td>
<td>10</td>
</tr>
<tr>
<td>Avg</td>
<td>7.6</td>
<td>7.1</td>
<td>74.7</td>
<td>13.3</td>
<td>1.7</td>
<td>0.5</td>
</tr>
<tr>
<td>Pärnu, carbonate</td>
<td>4 Min</td>
<td>25.2</td>
<td>15.5</td>
<td>75</td>
<td>15</td>
<td>0</td>
</tr>
<tr>
<td>Rezekne, carbonate</td>
<td>Max</td>
<td>75</td>
<td>43</td>
<td>85</td>
<td>25</td>
<td>0</td>
</tr>
<tr>
<td>Avg</td>
<td>45.5</td>
<td>25.1</td>
<td>80</td>
<td>20</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total samples</td>
<td>Max</td>
<td>75</td>
<td>69</td>
<td>99.8</td>
<td>46</td>
<td>25</td>
</tr>
<tr>
<td>Avg</td>
<td>23.9</td>
<td>22.4</td>
<td>76.2</td>
<td>18</td>
<td>1.3</td>
<td>0.6</td>
</tr>
</tbody>
</table>

The mineralogical composition of the heavy fraction was studied in 131 samples from four boreholes (Table 3, Fig. 8b). The heavy fraction accounted for 0–6.7% of siliciclastic (average 0.7%) and 0–64.1% of carbonate rocks (average 8.3%). The heavy fraction of carbonate rocks consisted of goethite and hematite (average 30.4%), transparent heavy minerals (22.9%), ilmenite and magnetite (21.6%), pyrite (14.1%), biotite (4.8%), muscovite (2.1%), barytes (1.8%), leucoxene (1.7%), and chlorite (0.6%) (Table 3, Fig. 8b). The heavy fraction of siliciclastic rocks included transparent heavy minerals (average 33.1%), biotite (22.1%), ilmenite and magnetite (19.9%), muscovite (4.2%), goethite and hematite (8.4%), leucoxene (5.4%), pyrite (3.2%), chlorite (1.8%), and barytes (1.7%). The biotite, chlorite, muscovite, and leucoxene contents were higher in the heavy fraction of siliciclastic rocks, while pyrite, goethite, and hematite showed higher values in carbonate rocks. The barite content was approximately the same, but the average ilmenite and
Fig. 8. Average mineral composition (in per cent) of carbonate and siliciclastic rocks in (a) clay fraction (Table 2) and (b) heavy fraction (Table 3). Dol, dolomite component; Cl, clay fraction; I, illite; Ch, chlorite; Kl, kaolinite; Mo, montmorillonite; MC, montmorillonite-chlorite; G, goethite; H, hematite; Q, quartz; F, feldspar; Si, siderite; HF, heavy fraction; Bi, biotite; Mu, muscovite; Ba, barytes; P, pyrite; Fe, Fe-oxides; L, leucoxene; IM, ilmenite and magnetite; TH, transparent heavy minerals.

Magnetite content was higher in carbonate rocks (Fig. 8b). Magnetite occurred more often in carbonate rocks, while ilmenite prevailed in siliciclastic rocks. Sometimes sulphides (pyrite and sphalerite) were recorded in the thin sections of dolostones (Fig. 5a,b) and dolomitic marlstones. Some siliciclastic samples included 1–2% goethite and hematite (Fig. 6a,b). Hematite, goethite, and pyrite could be identified in some thin sections of mixed carbonate-siliciclastic rocks (Fig. 6c,d).
Table 3. Heavy mineral fraction (0.05–0.1 mm) of the rocks of the Mehikoorma (421), Ruhnu (500), Tartu (453), and Valga (10) cores, determined by immersion mineralogical analysis

<table>
<thead>
<tr>
<th>Formation, rocks</th>
<th>Number of samples</th>
<th>Statistical parameter</th>
<th>Heavy fraction, %</th>
<th>Mineral component, %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Biotite</td>
<td>Chlorite</td>
</tr>
<tr>
<td>Aruküla, siliciclastic</td>
<td>51 Min</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Kernave, carbonate</td>
<td>Max</td>
<td>1.6</td>
<td>93.2</td>
<td>15.5</td>
</tr>
<tr>
<td>Kernave, siliciclastic</td>
<td>Avg</td>
<td>0.4</td>
<td>22.7</td>
<td>1.5</td>
</tr>
<tr>
<td>Kernave, carbonate</td>
<td>Min</td>
<td>0.4</td>
<td>1.0</td>
<td>0</td>
</tr>
<tr>
<td>Kernave, carbonate</td>
<td>Max</td>
<td>32.6</td>
<td>1.5</td>
<td>0.1</td>
</tr>
<tr>
<td>Kernave, carbonate</td>
<td>Avg</td>
<td>16.52</td>
<td>1.2</td>
<td>0</td>
</tr>
<tr>
<td>Leivu, siliciclastic</td>
<td>Min</td>
<td>0</td>
<td>4.6</td>
<td>0</td>
</tr>
<tr>
<td>Leivu, carbonate</td>
<td>Max</td>
<td>2.9</td>
<td>73.0</td>
<td>6.8</td>
</tr>
<tr>
<td>Leivu, carbonate</td>
<td>Avg</td>
<td>0.5</td>
<td>38.3</td>
<td>2.7</td>
</tr>
<tr>
<td>Vadja, carbonate</td>
<td>Max</td>
<td>64.1</td>
<td>34.7</td>
<td>3.6</td>
</tr>
<tr>
<td>Pärnu, siliciclastic</td>
<td>Min</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Pärnu, carbonate</td>
<td>Max</td>
<td>6.5</td>
<td>8.3</td>
<td>0.4</td>
</tr>
<tr>
<td>Pärnu, carbonate</td>
<td>Avg</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Pärnu, carbonate</td>
<td>Max</td>
<td>6.7</td>
<td>53.0</td>
<td>7.2</td>
</tr>
<tr>
<td>Pärnu, carbonate</td>
<td>Avg</td>
<td>1.3</td>
<td>5.3</td>
<td>1.2</td>
</tr>
<tr>
<td>Pärnu, carbonate</td>
<td>Min</td>
<td>0.4</td>
<td>0.1</td>
<td>0</td>
</tr>
<tr>
<td>Total samples</td>
<td>Min</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total samples</td>
<td>Max</td>
<td>64.1</td>
<td>93.2</td>
<td>15.5</td>
</tr>
<tr>
<td>Total samples</td>
<td>Avg</td>
<td>4.5</td>
<td>13.4</td>
<td>1.2</td>
</tr>
</tbody>
</table>

**INTERPRETATION AND DISCUSSION**

Primary variation in the total iron content of Palaeozoic sedimentary rocks has been explained by different influx of detrital grains and various weathering and erosion during transgressive and regressive sea-level changes and climate changes (Ellwood et al. 2000, 2001). During diagenesis the total iron content of sediments could increase, decrease or iron could change its valence (Elmore 1993; Brand 1994; Mücke 1994; Shogenova 1999; Vigliotti et al. 1999; Yamazaki et al. 2003; Funk et al. 2004; Zwing et al. 2005). These processes are mainly controlled by redox potentials of diagenetic fluids. Iron oxides dissolve in suboxic and anoxic (sulphate-reducing) environments, but precipitate in oxidation conditions (Passier et al. 2001).
The Fe$_2$O$_3$/Al$_2$O$_3$ ratio can serve as an indicator of detrital input of iron minerals versus diagenetic input. In this case Clark values of iron and aluminium and their relation in the specific rock type (Turekian & Wedepohl 1961) is used for calculating the excess of the element detrital level (Thomson et al. 1995). The Fe/Al$_{detrital}$ ratios for a global survey of carbonates by Turekian & Wedepohl (1961), which have been recalculated into oxides (Fe$_2$O$_3$/Al$_2$O$_3$$_{detrital}$), are 0.68 for carbonates and 0.3 for sandstones. These values coincide with the average values of the Fe$_2$O$_3$/Al$_2$O$_3$ ratio which we got for our Devonian dolostones and siliciclastic rocks (Table 1). Larger values reflect diagenetic mineralization, while smaller values than the Fe$_2$O$_3$/Al$_2$O$_3$$_{detrital}$ ratio are the result of diagenetic corrosion and dissolution. In the investigated Devonian rocks the total iron content is not very high and mainly correlates with the clay content (Fig. 4a–c). However, the Fe$_2$O$_3$/Al$_2$O$_3$ ratio shows the available excess of iron over detrital input in some carbonate samples, mainly from the Vadja Formation (Fig. 2, Table 1). The highest Fe$_2$O$_3$/Al$_2$O$_3$ ratio was obtained for dolostones of the Mehikoorma (421) (Fig. 5a) and Häädemeeste cores (Fig. 5b). Iron minerals in these rock samples, represented by Fe-oxides (hematite and goethite) and Fer-sulphides (pyrite and sphalerite), are obviously of diagenetic origin and could have formed in an oxidizing environment. In the other carbonate samples the Fe$_2$O$_3$/Al$_2$O$_3$ ratio is lower than the Fe$_2$O$_3$/Al$_2$O$_3$$_{detrital}$ ratio, which could be indicative of a reducing environment. Significant correlation of Fe$_2$O$_3$ with the Al$_2$O$_3$ content for all rock types suggests prevailing primary accumulation of iron-bearing minerals and possible changes in their form during diagenesis. Variations from the Fe$_2$O$_3$/Al$_2$O$_3$$_{detrital}$ ratios of 0.68 and 0.3 indicate diagenetic increase and decrease in the total iron content also in siliciclastic rocks.

As stated above, the total iron contents of red, pink, mottled, and grey rocks are similar. Such a range of the total iron content is in agreement with data from other “red beds” (Mücke 1994) and also proves that the colour of the rocks does not depend on the iron content, but is mainly determined by its form. Red coloration is mainly determined by hematite or, in other words, by excess ferric iron, in which Fe$_2$O$_3$/FeO ratios > 2 (Turner 1980). The formation of hematite can be explained by frequently changing redox conditions during diagenesis. A low water table and absence of organic matter are favourable for an oxidizing environment and formation of hematite during early diagenesis (Van Houten 1973). The FeO content of Estonian Devonian rocks is rather low – in a range of 0.03–0.43% in carbonate, 0.05–0.52% in siliciclastic, and 0.01–0.94% in mixed rocks. The average FeO content values in all rocks are between 0.16 and 0.24%, being the highest in marlstones and mixed rocks. In the Mehikoorma (421) core the Fe$_2$O$_3$/FeO ratio is rather high (Fig. 2), remaining below 2 only in a few sandstones. The highest ratio was observed in the mainly mottled and red rocks of $D_{3nrK}$–$D_{2atr}$, while the mainly grey succession of $D_{4rz}$–$D_{3nrV'}$ revealed a lower total iron content and a smaller Fe$_2$O$_3$/FeO ratio. It should be noted that such a low FeO content as in the Devonian has been recorded only in limestones and
argillaceous limestones of the Estonian Ordovician and Silurian sequences, but the average total iron content of the latter rocks is significantly lower (0.3–1.1%; Shogenova 1999). Ordovician–Silurian marlstones, dolomitic marlstones, and dolostones of different genesis have an average FeO content in the range of 0.29–1.02% and Fe$_2$O$_3$$_{total}$ in the range of 0.64–2.96%, with the highest values in glauconite-bearing carbonate rocks and secondary dolostones (Shogenova 1999). The average total iron content of the studied Devonian carbonates is 1.56–2.52%, indicating excess Fe$_2$O$_3$ in Devonian rocks in comparison with the other Palaeozoic rocks. The occurrence of Fe(III) may be evidence of a prevailing oxidizing environment during early diagenesis.

The studied carbonate rocks have a higher Fe$_2$O$_3$$_{total}$/Al$_2$O$_3$ ratio than siliciclastic rocks, while the siliciclastics have higher TiO$_2$/Al$_2$O$_3$ and K$_2$O/Al$_2$O$_3$ ratios (Table 1, Fig. 4a–c). According to Turekian & Wedepohl (1961), the global average values of the last two ratios are also higher in carbonate rocks than in sandstones. Our results of chemical analysis of the Devonian rocks are explained by (1) higher average goethite, hematite, magnetite, and pyrite contents in heavy fractions of carbonate rocks (Fig. 8b), (2) higher contents of leucoxene and ilmenite in siliciclastic rocks (Fig. 8b), and (3) transformation of minerals from the clay fraction into hematite during early diagenesis preceding dolomitization. The presence of pyrite, sphalerite, and siderite (in small amounts) in carbonate rocks is also supported by the absence of Fe$_2$O$_3$$_{total}$–FeO correlation in carbonate rocks (Fig. 4e). In contrast, the FeO content in a part of mixed and siliciclastic rocks correlates with the Fe$_2$O$_3$$_{total}$ content. This can be explained by the occurrence of Fe(II) in the clay fraction of mixed carbonate-siliciclastic and siliciclastic rocks. In the other samples FeO correlates with the TiO$_2$ content, indicating the occurrence of ilmenite in the heavy fraction. Some samples located lower on the Fe$_2$O$_3$$_{total}$–FeO plot than the correlated group (their Fe$_2$O$_3$$_{total}$/FeO proportion is higher than in other rocks) may also include pyrite and sphalerite in the heavy fraction. The negative correlation of the total iron content with the MgO content in carbonate rocks and the absence of any correlation between these parameters in siliciclastic rocks (Fig. 4d) suggests a decrease in the content of iron minerals during dolomitization of carbonate rocks and redistribution of iron minerals during dolomite cementation of siliciclastic rocks (Fig. 9). The first explanation could not be checked using comparison of Devonian limestones with dolostones due to complete dolomitization of the studied carbonate rocks. However, comparison of the Devonian dolostones with a total Fe content of 0.63–3.3% in the Mehikoorma (421) core with the underlying Upper Ordovician limestones shows a lower range of the total iron content for limestones (0.86–2.23%) (Shogenova et al. 2005). However, the Ordovician limestones have a higher average (1.76%) than the Devonian dolostones (1.56%). This also supports the conclusion about the general decrease in the total iron content of Devonian dolostones with a few cases of authigenic secondary mineralization where the Fe$_2$O$_3$$_{total}$/Al$_2$O$_3$ ratio is higher than the Fe$_2$O$_3$$_{total}$/Al$_2$O$_3$$_{detrital}$ ratio.
Iron-bearing minerals have altered significantly during diagenetic processes (Fig. 9; Klesment & Paap 1978; Klesment 1984, 1998). In early diagenesis dispersed fine grains and aggregates of pyrite precipitated in sediments as a result of the bacterial reduction process (Fig. 5a,b). Later, larger pyrite crystals formed in vugs, fractures, and sandstone matrix, prior to late diagenetic replacement of dolomite by calcite (Fig. 6d). The formation of mottled, patchily distributed iron oxides (Figs 5c,d; 6a–c) is presumably connected with the middle phase of diagenesis (Fig. 9). The red colouring of sandstones due to grain coatings may be of early diagenetic origin (Kuršs & Stinkule 1972), but some goethite and hematite could also have formed in later diagenetic stages from magnetite. Goethite and hematite in carbonate rocks could be of primary and diagenetic origin. Primary iron-bearing minerals were formed during sedimentation processes and their content correlates with the clay content (Figs 4a, 9). Secondary iron minerals do not correlate with the clay content as they formed from clay minerals and redistribution of iron is associated with dolomitization. Dolostone samples with secondary iron minerals show a poor correlation on the Fe$_2$O$_3$total–Al$_2$O$_3$ plot, and their Fe$_2$O$_3$total/Al$_2$O$_3$ ratio is higher than the Fe$_2$O$_3$total/Al$_2$O$_3$detrital ratio (Fig. 2). In contrast, some of the carbonate and siliciclastic samples show a low total iron content for a given Al$_2$O$_3$ content (Fig. 4a). This decrease in the iron content may
be connected with diagenetic processes during which some iron minerals could have dissolved in a reducing environment. In general, the total iron content decreased during dolomitization of carbonate rocks, but its concentration does not change significantly in siliciclastic rocks. Devonian siliciclastic rocks of Estonia have also a higher total iron content than Cambrian siliciclastics (Shogenova et al. 2001). Devonian dolostones have a similar, but on average lower total iron content than Upper Ordovician rocks and a significantly lower iron content than Middle Ordovician rocks, probably due to late-diagenetic origin of the (Middle) Ordovician dolostones of the Mehikoorma (421) core (Shogenova et al. 2005).

CONCLUSIONS

(1) The IR of the Devonian carbonate rocks of Estonia consists mainly of illite, illite-smectite, chlorite, kaolinite, quartz, and K-feldspar, with minor hematite, goethite, pyrite, magnetite, and siderite. The IR of siliciclastic rocks includes mainly quartz, K-feldspar, illite, and chlorite, with minor Ti-bearing minerals, hematite, goethite, siderite, and pyrite.

(2) The total iron content of the rocks was controlled mainly by detrital input during the sedimentation process. Iron-bearing minerals changed their form during diagenesis and Fe(III) iron minerals became dominant due to an oxygen-rich environment, low water table, and arid climate.

(3) Red coloration of siliciclastic rocks due to hematite coatings of quartz grains could be early diagenetic in origin, or could be also formed from magnetite and other iron oxides during later diagenetic stages. Red coloration of carbonate rocks (dolostones and marlstones) may by of early diagenetic origin before dolomitization, but iron pigmentation was redistributed also in the later diagenetic stages.

(4) The average values of the $\text{Fe}_2\text{O}_3/\alpha_\text{Al}_2\text{O}_3$ ratio coincide with the Fe/Al ratios in the sandstones and carbonates of the Earth’s crust. The TiO$_2$/Al$_2$O$_3$ and K$_2$O/Al$_2$O$_3$ ratios are higher in Devonian siliciclastic rocks of Estonia, but are higher in the carbonates of the Earth’s crust (Turekian & Wedepohl 1961).

(5) The iron oxides and hydroxides can be of primary, early diagenetic and/or late diagenetic (secondary) origin (Fig. 9). Detrital magnetite could have oxidized to hematite and goethite, while detrital ilmenite could transform to leucoxene during early diagenetic stages. Authigenic pyrite could have formed at all diagenetic stages. During dolomitization and dolomite cementation detrital iron minerals could be corroded and the total iron content could decrease from primary detrital values. Late diagenetic authigenic mineralization of sulphides associated with fractures and vugs took place in dolostones. Clay minerals could serve as a source of iron for authigenic mineralization leading to increase in total iron both in carbonate and siliciclastic rocks.
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**Diageneetiliste muutuste mõju rauamineraalidele Eesti Devoni karbonaatsetes ja terrigeensetes kivimites**

**Alla Šogenova ja Anne Kleesment**

vältel valitsenud oksüdeerivad tingimused, kuiv kliima ja madal veetase. Hema-
tiitse kattega kvartsterade massilise esinemise põhjustad terrigeensete kivimite punavärvilisus on tõenäoliselt varadiagenetiline, kuid neis kivimeis on toimusud ka hilisemal diageneesil hematiidi moodustumine magnetiidi arvel ja rauaoksüü-
dide pigmenti moodustumine karbonaatsetes kivimites. Dolokivide ja dolomii-
merglite punavärvilisus on valdavalt varadiagenetiline, eelnedes dolomitsatsiooni-
protsessidele, kuid mõningane värvuste muutumine võib olla seotud ka hilisdia-
geneseiga. Rauamineraalide tekke seostamine erinevate kivimi moodustumise staadiumidega (settimine, vara-, kesk- ja hilisdiagenesees) põhineb mineraloogilistel uuringutel immersioonis ja õhikutes, arvestades ka keemilise koosseisu andmeid. Keskmise Fe²O₃/Al₂O₃ suhe on karbonaatkivimites kõrgem kui terrigeensetes kivimites, mis on vastavuses ka maakoore kohta arvutatud andmetega. See näitab, et rauaühendite ümberjaotumine diageneesil toimub intensiivsemalt just terrigeens-
setes kivimites. Devoni terrigeensete kivimite Ti₂O/Al₂O₃ ja K₂O/Al₂O₃ suhete väärused on maakoore keskmistest mõnevõrra kõrgemad, mis võib osutada siin savikivimitse muutustele diageneesi käigus.