This paper presents data on the geochemical and mineralogical characteristics of a core pertinent to the limnological history of Lake Iidesjärvi in the town of Tampere (S. Finland). The geochronology together with the pollen and diatom stratigraphy of its sediment sequence have been given earlier (Alhonen 1981). The new palaeolimnological results from Lake Iidesjärvi show the anthropogenic enrichment of heavy metals such as Cd, Cu, Hg, Pb and Zn in the topmost sulphide-banded clay-gyttja of its lithostratigraphy. This cultural eutrophication was interpreted by studying extraction kinetics of Fe, Si, Ca, Mg, K, Cu and Zn using ammonium oxalate and hydroxylammonium chloride plus acetic acid solutions. The productivity history of Lake Iidesjärvi was studied by the determinations of ignition loss, sedimentary chlorophyll and sedimentary humus. The mineralogical composition of the lithostratigraphy shows that the presence of clay minerals, particularly vermiculite, with extractable Mg, Fe and Na has probably led to minero-eutrophy of Lake Iidesjärvi since its isolation from the Ancylus Lake in agreement with the results of the diatom stratigraphy.

Keywords: clay minerals, lithostratigraphy, geochemistry, sediments.

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Introduction

Earlier studies (Alhonen 1981) show that Lake Iidesjärvi in the town of Tampere (South Finland) has been limnologically more or less productive since its isolation from the Ancylus Lake. This conclusion is based on the diatom stratigraphy of the main core, which was taken with a Livingstone sampler from the deepest part of the lake (Fig. 1). Lake Iidesjärvi has also been affected by intensive land use and settlement of the town of Tampere. This is reflected in the lithostratigraphy as the forma-
tion of sulphide bands in the topmost clay-gyttja indicating the dynamics of cultural eutrophication (see e.g. Alhonen 1979). Limnochemically, oxygen deficiencies are therefore common because of the high nutrient level caused by sewage. The limnological status of Lake Iidesjärvi and of its geological environment has been reported earlier by Alhonen (1981, pp. 98–99).

In this report we present data on the mineral and geochemical characteristics of the bottom sediments pertinent to the limnological history of Lake Iidesjärvi. The purpose is thus to obtain information on the nature of the sedimentary material in relation to the geology of the drainage basin, the major source of the sedimented material being the drainage of the lake rather than the organic matter synthesized in its water (Mackereth 1966). The analyses were done on the core (Fig. 2) described by Alhonen in 1981 (p. 100 and Fig. 3). The uppermost 40 cm were analysed in greater detail, because they were deposited during the pollution development of Lake Iidesjärvi.

**Analytical methods**

**Geochemistry**

The homogenized subsamples were dried for 24 hours at 90°C for acid digestion and ignition loss determinations. The samples were air-dried at room temperature for extraction procedures. The loss on ignition was determined by igniting the samples for 1 hour at 800°C.
About 1 g of sample was weighed into a Teflon dish and 5 ml of HCl (12 M), 5 ml of HNO₃ (15.5 M) and 0.4 ml of HClO₄ (16.6 M) were added. The samples were digested on a warm plate at 90°C for 12 hours. The acids were then evaporated until HClO₄ vapours developed at 160°C. The samples were cooled, 2 ml of HCl (12 M) were added and the salts were dissolved by boiling for 5 minutes. The liquid was removed with pure water and transferred to flasks 10 ml in volume for analysis. A number of samples were extracted for organically bound elements with sodium hypochlorite solution, for sedimentary chlorophyll with acetone, and for humic substances with NaOH solution (Table 1). Elemental forms and binding of elements were studied by subjecting the samples to a selective sequential extraction procedure using ammonium acetate solution (NH₄Ac), solution of hydroxylammonium chloride (HXI), and a mixture of HXI-HAc. Solution kinetics were studied using NH₄Ac and HXI-HAc extraction and varying the time intervals (Table 1).

The acid-digested samples were analysed for Al, Ca, Cd, Cu, Fe, Hg, K, Li, Mg, Pb, Rb, Zn, PO₄ and sedimentary humus (Table 2). The NH₄Ac and HXI-HAc extract supernatants were centrifuged at different time intervals and analysed for Ca, Cu, Fe, K, Mg,
Table 2. Analytical procedures.

| CONSTITUENT DETERMINED | WAVELENGTH nm | SLIT | FLAME | METHOD |
|------------------------|---------------|------|-------|--------|
| Cd                     | 228.80        | 4    | air-C₂H₂ | AAS    |
| Cu                     | 324.75        | 4    | »      | »      |
| Fe                     | 248.82        | 3    | »      | »      |
| K*                    | 766.49        | 4    | »      | »      |
| Li*                    | 670.78        | 4    | »      | »      |
| Mn                     | 279.48        | 3    | »      | »      |
| Pb                     | 283.31        | 4    | »      | »      |
| Rb*                    | 780.02        | 5    | »      | »      |
| Zn                     | 213.86        | 4    | »      | »      |
| Ca*                    | 422.67        | 4    | N₂O-C₂H₂ | AAS    |
| Al*                    | 309.27-309.28 | 4    | »      | »      |
| Mg*                    | 285.21        | 4    | »      | »      |
| Hg                     | 253.60        | 4    | »      | AAS Cold vapour |
| Si                     | K** 769.90    | 4    | »      | Visible spectrophotometric |
| PO₄<sup>Sedimentary</sup> | Li 670.78    | 4    | »      | »      |
| PO₄<sup>Chlorophyll</sup> | Li 670.78   | 4    | »      | »      |
| PO₄<sup>Sedimentary humus</sup> | Sr 460.70 | 4    | »      | »      |

*) 1 % of NaCl added for ionization buffer  
**) hollow cathode lamp used

Mn, Zn and Si (see Table 2). The HX1 and sodium hypochlorite extracts were also analysed for metals. An appropriate matrix was used in standard solutions in the analysis of the extracts. The spectrophotometric determinations were accomplished by AAS. Si was determined as blue molybdosilicic acid using oxalic acid to eliminate PO₄ interference (Babko and Pilipenko 1976). PO₄ was determined as blue molydophosphoric complex using ferrous iron and sulphite as the reductant for iron (Babko and Pilipenko 1976). After reduction of iron the sedimentary humus was determined from an aliquot, the solution absorbance being measured at wavelength 460.7 nm (Table 2). The acetone extract absorbances were measured for sedimentary chlorophyll at wavelength 670.78 nm (Vallentyne 1955; Moss 1967 and Lorenzen 1967). The absorbances of the NaOH extract were measured after coagulating the cations and colloidal iron with NaSO₄ by a spectrophotometer (Hitachi, Model 101).

Some comments on geochemical methods

According to Agemian and Chau (1976) many of the trace metals in lake sediments can be extracted by acid digestion and it is assumed that the total leachable elements represent the most characteristic form of elements of interest when studying the sedimentary record for palaeolimnological evidence. The humic matter is generally extracted by NaOH solution, but an identical result is achieved using HCl extraction (Schnitzer and Skinner 1968). The HClO₄ used in the present study, however, caused partial oxidation and degradation of the
humic macromolecules (Maximov et al. 1977 and Ruggiero et al. 1980). Nevertheless, the results correlate with those obtained from NaOH extraction and seem to represent total humus.

Ammonium acetate extract was used to liberate the «exchangeable metals». Thermodynamic calculations show that NH₄Ac at pH 4.8 will not dissolve iron and manganese oxides, oxyhydroxides or hydroxy complexes (Smith and Martell 1976). Thus the elements released appear to be primarily specifically adsorbed, although Ca²⁺ will be complexed by the NH₄Ac extract (Sillen and Martell 1971). Carbonates dissolve, but silicates and organic C are not attacked. Heavy metals from labile organic complexes (e.g. humic acid complexes) are also dissolved to some extent by the NH₄Ac extract (Slavek et al. 1982). The hydroxylammonium chloride extract dissolves oxidized forms of Mn (Chao 1972) but does not attack the Fe oxyhydroxides and oxides. The 1 M hydroxylammonium chloride with 4 M acetic acid dissolves poorly crystallized Fe oxyhydroxides and oxides (Chester and Hughes 1967). It also attacks siliceous material, especially during prolonged extraction. Metals from labile organic complexes dissolve to some extent. Sodium hypochlorite at pH 9.3 liberates the heavy metals complexed by organic matter (Gibbs 1973). Analytical accuracy tends to be better than ± 5 per cent.

Mineralogy

For qualitative determination of the clay minerals and residual rock-forming minerals, 28 X-ray diffraction analyses were performed on the fine-grained (< 20 μ) material from different parts of the sedimentary column (see Fig. 2), using a Philips X-ray diffractometer with CuK radiation (cf. Uusinoka 1975, p. 13).

Results and discussion

The content of acid-digested elements, Ca, Li, Mg, Al and, especially, Fe, is elevated in the lowermost banded sulphide clay. As indicated by loss on ignition, sedimentary, chlorophyll and humus determinations, the productivity of Lake Iidesjärvi is initiated immediately after its isolation. With the exception of Fe, the content of elements increases, too (see Figs 3–6). Thereafter, as soon as the sedimentation of the brown detritus gyttja is interrupted by the pale grey gyttja layer (see Fig. 2) the organic matter decreases together with a temporary increase in the content of Li, Al, Mg, Ca, Zn and PO₄. First the K and Rb contents reach a minimum whereas the organic matter increases further to a maximum as the sedimentation of the brown fine detritus gyttja continues. At this point the Mn and Cu contents are elevated, too, and a slight decrease is shown in the Fe content. The organic matter then increases again together with the elevated content of most of the elements studied. At a depth of c. 270–230 cm, the organic content decreases once more. The Cu content correlates with the organic matter, showing, however, no depressions. The humus content (Fig. 4) shows a maximum at a depth of 170 cm as does the NAP curve (see Alhonen 1981). The Li, K and Rb contents are elevated at a depth of 220–190 cm, and the Mn content at a depth of 190 cm.

The Al, PO₄, Zn, Cu and Li contents increase in the topmost part of the lithostratigraphy. The elevated Zn and Cu contents seem to originate from anthropogenic enrichment whereas the Al content may reflect cultural acidification by SO₅ and NO₅ compounds emitted during fossil combustion in local industry or transported by long-range acid rain. As a consequence of acidification Al is mobilized and transported to the lakes
Fig. 3. Lake Iidesjärvi: vertical distribution of loss on ignition, sedimentary chlorophyll and Fe in the investigated core.

Fig. 4. Lake Iidesjärvi: vertical distribution of Cu, Zn, PO$_4$ and sedimentary humus in the investigated core.

(Cronan and Schofield 1979 and Johnson 1979).

Anthropogenic enrichment of heavy metals such as Cd, Cu, Hg, Pb and Zn is clearly seen in the uppermost part (from 20 to 1 cm) of the shorter core (Figs 7 and 8), which was sampled for analysis with a denser sample interval than the longer one. The contents of Cu, Pb and Zn during the pollution development of Lake Iidesjärvi coincide more closely with those from heavily industrialized areas of southern Lake Michigan (Leland et al. 1973) and Lake Erie (Nriagu et al. 1979) than with those from Kuopio (Salminen 1976 and 1979) and Quebec (Ouellet and Jones 1982). The primary cause of anthropogenic enrichment of Cd, Cu, Hg, Pb and Zn is industrial development around Tampere since the beginning of this century, especially since the 1950s.

The increase of Mn content in the topmost part of the core (Fig. 9) may be due to diffusion and oxidative or organic fixation of the Mn$^{2+}$ released after being reduced as a consequence of cultural eutrophication of Lake Iidesjärvi. It is also indicated by the increase of organic matter and PO$_4$ content in the uppermost lithostratigraphy as dis-
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Fig. 5. Lake Iidesjärvi: vertical distribution of Al, Ca, Mg and Mn in the investigated core.

Fig. 6. Lake Iidesjärvi: vertical distribution of Rb, K and Li in the investigated core.

cussed earlier by Alhonen (1979, pp. 21–22) and others. However, the increase of Mn is not so pronounced as in a Norwegian fjord discussed in a study by Skei and Paus (1979).

Some features caused by the cultural eutrophication of Lake Iidesjärvi can be discovered by extraction studies using ammonium acetate (NH$_4$Ac) and hydroxylammonium chloride plus acetic acid (HXl-HAc). The results show that the amount of loosely bound HXl-HAc-extractable Si is lower in the upper sediment sequence (12–100 cm) than in the sequence beneath it (110–170 cm). The same is shown by the HXl-HAc and NH$_4$Ac-extractable Fe, which correlates with Si (Fig. 10). During short extraction the HXl-HAc-soluble Si released is evidently loosely bound, but during longer extraction (from 40 to 640 minutes) Si is withdrawn from diatom cells (see Lewin 1961 and Jones and Handreck 1963) and clay minerals as well. The elevated Si solubility during 160–320 minutes’ extraction coincides with the elevated Zn solubility, which indicates that the solubilized elements originate primarily from clay minerals (Figs 11–14). The correlation between HCl-HAc-soluble Si, Fe and Ca indicates that
these elements interact chemically in the sediments of Lake Iidesjärvi (cf. Morgan and Stumm 1963). The amount of NH$_4$Ac-soluble Si is low and its variation great, which indicates weak adsorption of Si in the samples. In contrast, Si is strongly associated with the surface layer owing to freshly precipitated natural oxyhydroxide precipitates, and Ca, Mg, Mn and Zn are adsorbed (Vuorinen and Carlson 1983).

The depletion of loosely bound Si in the upper sediment sequence might thus be due to increased diatom uptake of dissolved Si from lake water as a result of cultural eutrophication caused by anthropogenic input of PO$_4$ (Powers and Ayers 1967; Schelske and Stroermer 1971). The fluctuation in K, Mg, Ca and Fe solubilities seems to reflect the pollution development of Lake Iidesjärvi in its sediment sequence (Figs. 15–18). As a rule, the amount of NH$_4$Ac-extractable elements considerably decreases during extraction, which is considered to indicate that the elements primarily exist as specifically adsorbed on the particulate material.

Fig. 7. Lake Iidesjärvi: vertical distribution of Zn, Cu and Cd in the topmost 40 cm of the investigated core.

Fig. 8. Lake Iidesjärvi: vertical distribution of Pb and Hg in the topmost 40 cm of the investigated core.
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Fig. 9. Lake Iidesjärvi: vertical distribution of loss on ignition Fe, Ca, P0\textsubscript{4} and Mn in the topmost 40 cm of the investigated core.

The adsorption order of the elements is calculated as the percentage extracted by NH\textsubscript{4}Ac solution during the shortest extraction time (15 min) of the total amount extractable, using average elemental contents of sample-intervals of 12 to 170 cm. The order was Mg, Ca, Mn, K, Zn, Cu (Fig. 19). This corresponds to the prevailing forms of loosely bound elements. Mg and Ca exist as cationic adsorbed forms as, to a slight lesser extent, do Mn, K and Zn. In contrast, some of the Fe was present in colloidal form in the sediments and oxidized partly to oxyhydroxides during the preparation of

Fig. 10. Lake Iidesjärvi: Fe and Si-HXl-HAc and NH\textsubscript{4}Ac dissolution (% of total soluble) from upper and lower parts of the investigated core during different extraction intervals.
Fig. 11. Lake Iidesjärvi: HXI-HAc-extractable Si in the investigated core during different extraction intervals.

Fig. 12. Lake Iidesjärvi: HXI-HAc-extractable Fe in the investigated core during different extraction intervals.

Fig. 13. Lake Iidesjärvi: HXI-HAc-extractable Ca in the investigated core during different extraction intervals.
Fig. 14. Lake Iidesjärvi: HXl-HAc and NH₄Ac-extractable Zn in the investigated core during different extraction intervals.

Fig. 15. Lake Iidesjärvi: NH₄Ac-extractable Fe in the investigated core during different extraction intervals.

Fig. 16. Lake Iidesjärvi: NH₄Ac-extractable K and Mg in the investigated core during different extraction intervals.
the samples. Cu associates mainly with organic matter (see also Presley et al. 1972; Tessier, Cambell and Bisson 1979).

To elucidate the character of the humic substances in the lithostratigraphy of Lake Iidesjärvi the absorbance of NaOH extracts of humic matter diluted 10 fold were measured at wavelengths of 430, 465, 490, 550, 600 and 665 nm, and absorption curves versus wavelength were constructed (Fig. 20). The slope of the curves is related to the condensa-
tion degree of the humic acids (Pauli 1967). A low slope is characteristic of highly condensed humic acids and a steep one of slightly condensed fulvic acids. The study showed that the fulvic acid component and the humus content increases with sediment depth, possibly because of the degradation of humic macromolecules in the sedimentary environment, e.g. by micro-organisms (see Anderson 1979). It is a subject worth studying further.

The relative order of elements solubilized by different extracts was calculated to evaluate the binding of elements by different sediment constituents. The element content digested in acid was taken as the total content digested in acid was taken as the total content. The appropriate percentage subdivision was calculated for each element after subtracting the other extractable element contents (Fig. 21). The following order was obtained:

| Elements bound by | extracted by | Order of binding |
|-------------------|--------------|-----------------|
| Silicate-clay matter acid digestion | HXl-HAc extraction | Zn Fe K Mg Mn Ca Cu |
| Fe oxyhydroxides and oxides | HXl extraction | Cu Fe Zn Ca Mn |
| Mn oxides | NH4Ac extraction | Mn Ca Zn |
| Adsorption | HClO extraction | Ca K Mg Cu Mn Zn Fe |
| Organic matter | | Cu Mn Zn |

In general, the mineralogical composition of the sediments was not dissimilar to that of the Finnish argillaceous sediments as a whole (see, e.g. Soveri 1956; Sippola 1974; Gardemeister 1975 and Alhonen et al. 1978). As to the residual, fine-grained and unweathered rock-forming minerals, quartz, plagioclase, potassic feldspar and amphibole were present in all samples throughout the

![Absorbance graph](image-url)
sedimentary sequence of Lake Iidesjärvi, with quartz being the dominant mineral and amphibole and potassic feldspar the least abundant. Mica and illite, in most samples both di and triocahedral types, and chlorite were present in varying amounts in all the samples. Vermiculite, too, was found throughout almost the whole lithostratigraphy of the lake as were the mixed-layer minerals chlorite-illite and illite-vermiculite. Identifiable amounts of smectite or kaolinite were not found in the samples analysed.

The presence of illite, chlorite and vermiculite, both as such and as members of mixed-layer minerals, indicates some progress of weathering phenomena before and during sedimentation of the sedimentary material derived from glacial drift eroded from bedrock situated, according to the orientation of the glacial striae, in an exceptionally wide sector, from Lake Iidesjärvi to N-SW (see Virkkala 1962). The composition of the bedrock varies conspicuously, the granitoids and various micaceous schists and gneisses, however, being dominant (see Matisto 1961).

The amount of clay minerals is closely correlated with the amount of exchangeable ions, such as potassium and magnesium, in soils. Owing to its high cation exchange capacity, vermiculite is particularly closely correlated with the contents of exchangeable cations (see e.g. Sippola 1974). In a sedimentation basin, the presence of clay minerals with extractable magnesium, iron and potassium might thus be able to create conditions leading to minero-eutrophy of a lake. Vermiculite has been found in the bottom deposits of Lake Iidesjärvi, since the lower part of the column studied, and its presence both as such and as a member of mixed layer illite-vermiculite, is somewhat higher than in most other areas studied (cf. Gardemeister 1975 and Alhonen et al. 1978). This is probably due to the micarich bedrock in the environment and the relatively easy wheatherability of biotite to the »hydromica» stage, i.e. biotite-vermiculite.

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