The distribution of porphyrins with different tumour localising ability among human plasma proteins

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Summary The distribution among the main fractions of human plasma lipoproteins of a number of porphyrins with different tumour localising ability has been determined by means of ultracentrifugation. A main trend is that the fraction of the dyes that are bound to low density lipoprotein (LDL) increases, and the fraction bound to HSA decreases with decreasing polarity of the dyes. An asymmetric charge distribution, such as that of TPPS2+ favours LDL-binding more than expected on the basis of lipophilicity. No correlation between the known tumour localising ability of the drugs tested in the present work and their relative affinity for LDL was found. One of the best tumour localisers reported in the literature, TPPS6, hardly binds to LDL, while Hp and Pp, which are commonly considered inefficient tumour localisers, do have a significant affinity for LDL. On the other hand, the LDL binding capacity for a drug is suggested to be a good index for cellular uptake. Such an index does not necessarily imply that the actual uptake occurs by the LDL pathway.

Materials and methods

Materials

Hp, protoporphyrin (Pp) and photoporphyrin (Ppp) were bought from Porphyrin Products (Logan, UT, USA). PhotofrinII (PII) (2.5 mg ml⁻¹) was obtained from Photomedica (Raritan, NJ, USA) and stored frozen in small vials. The tetraphenyl porphine sulphonates (TPPS₂₋₆) were kind gifts from Dr Bruce Burnham at Porphyrin Products. These sulphonated porphyrins belong to the same batch as that used by Kessel et al. (1987), and were tested chromatographically by that group. Their chromatographic analysis agreed with our own (Figure 4). Stock solutions of Hp, PII, TPPS₆ and TPPS₂ were made up in 0.03 M NaOH + 0.15 M NaCl, whereas dimethyl sulfoxide (DMSO) was stock solvent for Pp, Ppp, TPPS₂₋₆ and TPPS₂₋₆.

All other chemicals used were of the highest purity commercially available.

Fresh human plasma sampled in EDTA tubes was used in all experiments and was obtained from the same healthy person who had fasted overnight. All solutions for ultracentrifugation were made up with 0.1 M Tris-HCl buffer, pH 7.35, containing 0.4 mg EDTA (potassium salt) per ml.

Ultracentrifugation

A Beckman L8-70M ultracentrifuge with a fixed angle 70.1 Ti rotor was used. It was found that a 15 h run at 70,000 r.p.m. was suitable for a good separation of the different protein fractions. A 74 h run at 32,000 r.p.m., using a centrifuge with an SW-40 swinging bucket rotor, gave similar results, the only significant difference being a slightly better separation at the top of the gradient (i.e. in the VLDL/LDL region). The high-speed/short time method was used in this study, to reduce ultracentrifuge time and to reduce possible ageing effects on the samples. Our gradient, as well as those used by others (Chapman et al., 1981; Kelly & Kruksy, 1986), is slightly modified from that used by Redgrave et al. (1975). The top fraction of the gradient was made up using 0.15 M NaCl, but at higher densities we employed CsCl instead of NaCl/KBr, as CsCl gives a lower molarity of salt at a given density. This fact may be
important for the separation since it is well known that an increase in the salt concentration results in a decrease in porphyrin solubility. Furthermore, for the same reason we chose to apply the drug-containing plasma at the top of the gradient instead of at $d = 1.15 - 1.312 \text{g ml}^{-1}$ as employed by other groups (Chapman et al., 1981; Nilsson et al., 1981; Kelly & Krusky, 1986; Redgrave et al., 1975). Pure plasma gives a similar protein distribution pattern whether it was applied at the top or at the bottom of the gradient. Finally, we chose to use a density of 1.35 g ml$^{-1}$ at the bottom of the tube (Figure 1), instead of 1.21, to prevent the heaviest proteins and/or protein–porphyrin complexes from precipitating at the bottom. Several gradients known from the literature (Chapman et al., 1981; Kelly & Krusky, 1986; Nilsson et al., 1981) were tested in a series of experiments using the SW-40 rotor and found to give similar relative separations as the gradient used by us. A separate paper, giving the explicit experimental data on which our choice of method is based, will be published later.

Concentration
The porphyrin binding patterns obtained here refer to 7 µg ml$^{-1}$ plasma in the case of PII and 14 µg ml$^{-1}$ plasma in the case of all other porphyrins. Reducing the plasma concentration by a factor of 5 while keeping the total porphyrin concentration constant (i.e. 35–70 µg porphyrin per ml plasma equivalent to 2.5–5 mg per kg bodyweight) gave similar results as for 7–14 µg ml$^{-1}$ for all substances tested in this respect, i.e. Hp, PII, Pp and TPPS$_2$.

Analysis
Gradients are often analysed from the bottom up. It was found that this method resulted in a contamination of the HDL fraction by other heavy proteins and an apparatus was therefore constructed which analysed and fractionated the gradient from the top down. By means of a peristaltic pump and an LKB 2098 Uvicord III monitor with appropriate optical filters the absorbences (1 cm) at 276 nm (proteins) and at 405 nm (porphyrins) were continuously monitored. Fractions were also collected and measured spectrophotometrically (Cary 118) since the tetraphenyl porphine sulphonates have their absorption maximum at 417 nm in our solvent. From one sample to another the position of the protein maxima in the gradient might change slightly (i.e. by ± one fraction). However, with three exceptions (see Results) the maxima of the porphyrin distribution coincided exactly with the maxima of the protein distribution in the same run.

HPLC
High performance liquid chromatography was carried out with an RP18 column and a methanol/water gradient buffered at pH 7.4 as described previously (Sommer et al., 1984).

Results
The initial gradient and that obtained after centrifugation are shown in Figure 1. The protein distribution (Area$_{280\text{nm}}$), is shown in the upper parts of Figures 2 and 3. LDL, lipoprotein(a) (Lp(a)), HDL and heavy proteins (mostly human serum albumin (HSA)) are well separated. The separation is similar to the separations obtained by others by the use of lower speeds and longer times of centrifugation, except that in our high r.p.m. experiments VLDL is not resolved from the LDL peak. Using the present gradient, but lower centrifugation rates (SW-40 rotor) and longer times, the VLDL was separated from the LDL peak (data not shown). From such data it may be inferred that 5–15% of the LDL-area in Figures 2 and 3 is due to VLDL. The reproducibility from one run to another and between blood samples was generally better than 5% (data not shown). In our case, the reproducibility between separate runs was better than 10% of the values shown in the LDL region, the errors are larger (20%) in the case of low LDL binding.

Figure 1  Outline of the gradient before (left) and after centrifugation (15 h, 70,000 r.p.m.). The classification of the lipoprotein is according to Patch & Patch (1986).

Figure 2  (a) Protein distribution in the gradient. Mean value and standard errors from six individual samples. (b) Distribution in the gradient of the following porphyrins: TPPS$_3$ (○), TPPS$_{2\alpha}$ (△), TPPS$_{2\alpha}$ (△), TPPS$_{4\alpha}$ (□) and TPPS$_2$ (×). Reproducibility between separate runs better than 10% of the values shown (in the LDL region, the errors are larger (20%) in the case of low LDL binding).
Table 1 Distribution of porphyrins among human plasma proteins (centrifugation 15 h, 15°C, 70,000 r.p.m.)

| Porphyrin | HPLC ret. time (min) on RP18 | LDL | HDL | Heavy proteins |
|-----------|-------------------------------|-----|-----|----------------|
| Ppp       | 9.5                           | 10  | 57  | 34             |
| Hp        | 3.6, 4.0                      | 16  | 70  | 14             |
| PII       | 3.6–20                        | 22  | 41  | 37             |
| Pp        | 18                            | 1–2 | 18  | 80             |
| TPPS2a    | 0.05                          | 6   | 26  | 73             |
| TPPS2b    | 0.35                          | 7   | 17  | 38             |
| TPPS2c    | 3.95                          | 36  | 55  | 53             |
| TPPS2d    | 10.1                          | 60  | 80  | 10             |
| TPPS3     | 20.0                          |     |     |                |

Data from Reyfman et al. (1984) for Hp and Pp and Kessel et al. (1987) for the TPPS-series are included in parentheses for comparison.

Reproducibility of percentages bound to the different proteins better than 15% of the values shown, cf. legend of Figure 2.

Samples taken on different days was good, consistent with the s.e. bars shown on Figure 2. The maxima of the LDL, Lp(a), HDL and HSA peaks were found at fractions 4–5, 11–13, 17–19 and 28–29, respectively (Figures 2 and 3). In the cases of Hp, Pp, PII, TPPS, TPPS, and TPPS, (Figures 1 and 2), the maxima of the dye distribution always coincide with the maxima of the protein peaks. However, in the case of TPPS, the 417 nm peak in the HDL-region was shifted by one fraction to the right compared to the 280 nm peak and in the case of TPPS, the 417 nm peak in the HDL-region was shifted by three fractions to the right compared to the 280 nm peak. The latter shift is seen in the lower part of Figure 2. Furthermore, the major peak of the TPPS, adduct(s) is slightly shifted to the denser region of the heavy proteins by about one fraction, as seen in Figure 2.

Integration over distributions, as shown in Figures 2 and 3 by weighing the areas under the curves between given fractions (see legend to Figure 3), resulted in the data shown in Table I. Thus, in the present work we have not attempted to measure thermodynamic affinities but only the relative distributions of the drugs between the plasma proteins. We emphasize that the concentration of each plasma protein in the samples has remained constant to within 5% throughout the work. There may be a slightly different distribution of aggregated and monomeric species among the different serum proteins. However, absorbance measurements as performed in the present work appear to give correct distributions since fluorescence measurement was found to give similar distributions (data not shown). (The fluorescence quantum yields of porphyrins are very sensitive to the state of aggregation.) The porphyrin amount bound to VLDL was small compared to that bound to LDL (less than 15%) and therefore these two fractions are taken together. Only Pp and TPPS showed a tendency to bind to Lp(a). Even in these two cases, the fraction of the porphyrins bound to Lp(a) is small compared to that bound to LDL (<5%) and these two fractions are taken together in Table I.

When PII, Hp or Pp were incubated with plasma for different times (0, 4 and 16 h) at 37°C their distributions were almost identical to those shown in Figure 3.

Discussion

The relative binding of the porphyrins to LDL increases generally, as expected, with increasing lipophilicity (Table 1, Figure 3). Thus, Pp is more extensively bound to LDL than the structurally similar, but more polar Hp. Correspondingly, the relative binding of the porphyrins to the heavy proteins (mainly HSA) generally decreases with increasing lipophilicity, as best illustrated by the TPPS-series (Table I, Figure 3). However, there are important exceptions to this general trend: Ppp and Hp bind similarly in the heavy

Figure 3 (a) Protein distribution in the gradient. (b) Distribution in the gradient of the following porphyrins: Pp (△), Hp (○) and Ppp (□). (c) Distribution of PII in the gradient: Centrifugation: 15 h, 70,000 r.p.m.

Figure 4 (a) HPLC of a mixture of TPPS, TPPS, TPPS, and TPPS on a RP18 column eluted with a water/methanol gradient. The porphyrins were also run separately for identification of the peaks. (b) Distribution of the same porphyrins between LDL, HDL and heavy proteins (HSA on the figure). The percentage bound to each fraction was determined by weighing of distributions such as those shown in Figure 2. LDL, fractions 1–11; HDL, fractions 11–23; HSA, fractions 23–32.
protein region in spite of the fact that the latter porphyrin is significantly more polar than the former one. Similarly TPPS₄ binds more extensively to LDL than does TPPS₁, which is significantly less polar. This may be related to the asymmetric charge distribution on TPPS₂, which may cause a high affinity for a lipid/water interface. The asymmetry of TPPS₂₄ has been previously invoked by Kessel et al. (1987) as an explanation for their observation that TPPS₂₄ has a higher uptake in cells than TPPS₁₁. In summary, the relative binding capacity LDL has for a drug is mainly related to the lipophilicity of the drug, although other factors, such as properties of different sidegroups and the asymmetry/symmetry of the charge distribution play important roles for this affinity.

For most of the drugs we have tested, where these can be compared with results in the literature, there is generally broad agreement (Table I). However, in the cases of Hp and TPPS₂, we find relatively less in the heavy fraction and in the LDL fraction and more in the HDL fraction than did Reyftman et al. (1984) and Kessel et al. (1987). The discrepancy is even greater in the case of TPPS₄₄, for which we find much more in the HDL fraction and much less in the heavy protein fraction than did Kessel et al. (1987). The porphyrins which we find mainly in the heavier part of HDL, i.e. in HDL₄ (see Figure 1), seem in Kessel's experiments to have been less well resolved from the albumin peak than in our case. This may be related to an effect of salt on the relative binding of TPPS₄₄ as in his experiments that the porphyrin-labelled plasma was applied at a higher salt concentration. Indeed, we have shown (unpublished results) that, in the case of Hp, high salt causes more Hp to bind in the heavy pattern region. The fact that TPPS₂₄ and TPPS₄₄ seem to be bound more substantially to the heavier subfraction of HDL than the other porphyrins tested should be noted. These compounds are more polar than TPPS₂₄ and TPPS₄₄, just as the heavier subfraction of HDL is more polar than its lighter fraction. It is also noteworthy that the TPPS₄₄ peak at high densities seems to occur at a somewhat higher density than the protein peak in that region, implying that TPPS₄₄ appears to be bound not only to albumin but also to one or more of the other heavy proteins.

HpD injected in mice is reported to be lost much faster from the LDL fraction than from the HDL fraction (Kessel, 1986), as is Hp injected in mice and rabbits (Barel et al., 1986). Consequently, these observations would seem to indicate that the binding and release of these drugs to the lipoproteins is slow and that LDL is degraded faster than HDL. The same group showed for systemic administration of Hp to cancer patients that, although Hp is lost with similar rates from LDL and HDL, it is lost at lower rates from VLDL (Jori et al., 1984). However, from the present results the distributions of PII as well as those of Hp and Pp between the plasma proteins are similar for different incubation times ranging from 0 to 4 h (PII) and 0 to 16 h (Hp and Pp) at 37°C (data not shown). Thus equilibrium seems to be rapidly reached. Similarly, recent results obtained by ultracentrifugation of sera from patients injected with PII indicate that there is no change in the distribution of the drug with time between injection and sampling (Brown et al., work in progress), which is in agreement with our present results of rapid establishment of equilibrium. The possibility that there is a species difference in the relative stability of circulating porphyrin–lipoprotein complexes should be noted (but see below).

The assumption to be found in the literature that the tumour localising ability of porphyrins used in PDT may be related to their relative binding to LDL (Jori et al., 1984; Reyftman et al., 1984; Barel et al., 1986; Kessel, 1986) is not fully supported by the present work. Hp has a higher relative affinity for LDL than TPPS₁₁ and Pp has an even higher affinity (Table I), but Hp and Pp are generally considered inefficient tumour localisers. PII has a relative affinity for LDL lying between that of Hp and that of Pp but is a broad tumour localiser as reported by a number of authors (see the review by Moan, 1986). Moreover, TPPS₄ has a very low affinity for LDL (see Table I) and a relatively high affinity for heavy proteins, but is still one of the best tumour localisers studied so far, with respect both to absolute tumour uptake and to selectivity (Winkelman, 1985; Evenson, 1985; Peng et al., 1987). According to Kessel et al. (1987), albumin-bound drugs accumulate preferentially in stromal elements in tumour tissue while lipoprotein-bound drugs mediate intracellular localisation. As noted above, TPPS₄ appears to be bound not only to albumin but also to other heavy proteins which may contribute to its distribution in vivo.

The tumour selectivity of a drug may be related as much to the retention in tumour tissue as to its initial deposition. For this initial deposition, LDL transport may play an important role. Thus, it has been shown that a lipophilic fluorophore carried by LDL is selectively deposited in endothelial cells in the vasculature (Netland et al., 1985). Furthermore, strong PDT-effects on endothelial cells have been reported (Chaudhuri et al., 1987). The difference in retention of different porphyrins in cells is demonstrated by our HPLC experiments, showing that washing of HpD-loaded cells with a medium containing serum results in a selective removal of the monomeric components Hp, Pp and hydroxyethylvinyl deuteroporphyrin (Moan & Sommer, 1983).

Quite apart from the actual mechanisms of the drug uptake, there exists the possibility that the binding capacity for a drug (rather than its lipophilicity) is a good index for cellular uptake, as suggested by previous results for cellular uptake of TPPS analogues (Kessel et al., 1987).

We are forced to conclude that at present the determinants of tumour localisation of different drugs are poorly understood. LDL-binding may play one role, heavy proteins another. Other lipoproteins, such as VLDL, HDL (HDL₄ and/or HDL₃) and perhaps VHDL (very high density lipoproteins) as well as other factors such as aggregation properties, change in polarity with pH (Moan et al., 1980) chemical nature of side groups or the presence of different metal ligands (Hambrigt et al., 1975; Winkelman, 1967) may also be important.

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