Splenic eumelanin differs from hair eumelanin in C57BL/6 miceø

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The presence of melanin in spleens of black C57BL/6 mice has been known for long. Although its origin and biological functions are still obscure, the relation of splenic melanin to the hair follicle and skin pigmentation was suggested. Here, we demonstrated using for the first time electron paramagnetic resonance spectroscopy that black-spotted C57BL/6 spleens contain eumelanin. Its presence here is a “yes or no” phenomenon, as even in the groups which revealed the highest percentage of spots single organs completely devoid of the pigment were found. Percentage of the spotted spleens decreased, however, with the progress of telogen after spontaneously-induced hair growth. The paramagnetic properties of the spleen eumelanin differed from the hair shaft or anagen VI skin melanin. The splenic melanin revealed narrower signal, and its microwave power saturability betrayed more heterogenous population of paramagnetic centres than in the skin or hair shaft pigment. Interestingly, the pigment of dry hair shafts and of the wet tissue of depilated anagen VI skin revealed almost identical properties. The properties of splenic melanin better resembled the synthetic dopa melanin (water suspension, and to a lesser degree – powder sample) than the skin/hair melanin. All these findings may indicate a limited degradation of splenic melanin as compared to the skin/hair pigment. The splenic eumelanin may at least in part originate from the skin melanin phagocyted in catagen by the Langerhans cells or macrophages and transported to the organ.

Keywords: EPR, hair cycle, hair follicle, pigmentation, melanin degradation, spin-lattice relaxation

Melanin pigmentation is defined not only by melanogenesis but also by processing and transport of melanin granules from melanocytes to other cell types including but not limited to epidermal and hair follicle keratinocytes (Slominski et al., 2004; 2005). Forty years ago spleen of C57BL/6 mice was suggested to contain melanin (Weissman, 1967). This hypothesis was substantiated by histological demonstration of a black pigment in selected areas of murine spleen that was negative on histochemical iron (Pearls’ Prussian Blue) and lipofuscin (Ziehl-Neelsen) tests but positive on bleeching with potassium permanganate + oxalic acid (Weissman, 1967; van der Heijden et al., 1995). Ultrastructural studies also demonstrated that the majority of pigmented spleen cells contained melanosomes with fibrillar matrix, and therefore were defined as melanocytes (van der Heijden et al., 1995). Since normal spleen is a rather unusual site for the presence of melanin in the mammalian organism, it is mandatory to define the nature of the cells responsible for the collection, transport, accumulation and/or synthesis of this pigment in spleen. It is also important to define any correlations between splenic melanization and other physiological processes.

The presence of melanin-producing cells outside the skin has been demonstrated primarily in various organs of lower vertebrates (fish, amphibia and reptilia) where this pigment could be found in melanomacrophages — cells of a histiocytic character (Cicero et al., 1982; 1989; Sichel, 1988; Zuasti et al., 1989; Tsuji & Seno, 1990; Nakamura et al., 1993). These cells are responsible for the removal of pathogens and by-products of oxygen metabo-
lism, as well as for the re-utilization of iron in the hemopoietic system (Sichel, 1988; Zuasti et al., 1989; Tsuji & Seno, 1990; Nakamura et al., 1993). Thus, it was proposed (van der Heijden et al., 1995) that the murine splenic melanin is just an atavism of no systemic implications.

Van der Heijden et al. (1995) suggested that splenic melanin is a manifestation of cutaneous pigmentation of mice. Indeed, Weissmann (1967) did not observe any splenic pigmentation in albino mice. Therefore, we decided to focus our study on melanin deposition in the spleen in relation to the hair cycle-dependent melanogenesis in C57BL/6 mice. The C57BL/6 mouse is an excellent model to study hair cycle, its physiology and pathology as well as systemic immune functions (Paus & Cotsarelis, 1999; Stenn & Paus, 2001; Mueller-Roever et al., 2001; Slominski et al., 2004; 2005). The process of hair shaft pigmentation is strictly coupled to the hair cycle (Slominski et al., 1991; Slominski & Paus, 1993), a periodical turnover of cells which in anagen — the first phase of the cycle — create the pigmented hair shaft, and later undergo apoptosis in the next stage — catagen (Slominski et al., 2004; 2005). Also follicular melanocytes have been shown to undergo apoptosis at this stage (Tobin, 1998; Tobin et al., 1999). They may be replaced in the new hair cycle by a new pool of cells arisen from the appropriate progenitors (Nishimura et al., 2002), whereas the remaining apoptotic bodies which may contain melanin granules are phagocyted by Langerhans cells (Tobin, 1998; Slominski et al., 2004; 2005). The fate of this melanin is the most enigmatic process of hair pigmentation and melanocyte biology (Slominski et al., 1994; Tobin et al., 1999; Borovansky & Elleder, 2003; Sulaimon & Kitchell, 2003). Analogically to the case of lower vertebrates, one may hypothesize on the presence of melanophages which may transport some of the skin melanin to the place of its storage and further utilization. Melanin may be partially degraded in this process, because of its oxidative degradation in phagosomes of macrophages with the contribution of NADPH-oxidase, as suggested by Borovansky and Elleder (2003). Another possibility is deposition and degradation of melanin in organs destined to detoxify and reutilize blood, such as spleen or liver. Finally, melanin may remain in the cells of origin, but even there its further degradation cannot be excluded (Stanka et al., 1988). In contrast, melanized hair coat is periodically or continuously shed, while in epidermis melanin is degraded to form “melanin dust” and is removed together with dead cells (Borovansky & Elleder, 2003; Sulaimon & Kitchell, 2003; Slominski et al., 2004; 2005).

It must also be noted that the potential of various leukocytes, including neutrophiles, macrophages and lymphocytes to serve as a system of melanin transport in vertebrates, being an alterna-

tive or a supplement to the system of melanin transport within the “epidermal melanin units” was proposed already in the nineteenth century. In 1967 this idea was formulated by Wassermann, who also indirectly implied a similar notion concerning hair follicular pigmentation, and who connected both ideas with the phenomenon of visceral melanin deposition (Wasserman, 1967).

Hair cycling, including catagen and degradation of follicular melanocytes, is an immunologically-driven process (Paus & Cotsarelis, 1999; Stenn & Paus, 2001), which has systemic effects including, e.g., changes in the tempo of splenocyte proliferation (Slominski et al., 1997). Besides a direct engagement in the removal of follicular melanin phagocyted by the Langerhans cells in catagen, the spleen may participate indirectly in the cycle-related melanogenesis, or just passively reflect its activity. To address these questions we decided to focus on selected biophysical properties of skin and spleen melanin, in particular on their paramagnetic properties investigated by electron paramagnetic resonance (EPR) spectroscopy, being one of the most prominent, versatile and useful methods to investigate melanin in biological systems (Sarna & Plonka, 2005; Slominski et al., 2005). Therefore, the following questions were addressed with the help of EPR spectroscopy: 1. Is the presence of the black pigment in spleen a rule, a tendency or perhaps a property only quantitatively correlated with the stage of animal development/hair cycling stage? 2. Do the paramagnetic properties of the splenic pigment confirm its qualification as melanin, and what kind of melanin does it represent? 3. Is there a similarity between the splenic and hair/skin melanin? As the actual content and intensity of melanin production in skin changes in the telogen-anagen-catagen transition (Slominski & Paus, 1993; Slominski et al., 2005), which may additionally interfere with the corresponding phenomena in the spleen, we have chosen to focus in this pilot study on telogen — the “resting” phase of hair cycle, when there is no melanogenesis in hair follicles (Slominski et al., 1991).

**MATERIAL AND METHODS**

**Reagents and media.** Phosphate-buffered saline (PBS) without calcium and magnesium was purchased at Wytwórnia Surowic i Szczepionek BIOMED (Lublin, Poland). Ketamine for narcosis (Ketanest®) — at ASTA Medica AG (Frankfurt, Germany), beeswax, gum rosin, and 3,4-dihydroxy D,L-phenylalanine (D,L-dopa) — at Sigma-Aldrich (St. Louis, MO, USA).

**Animals.** Female inbred C57BL/6 mice 6–8-week-old (Animal Breeding Facility, Silesian Medical Academy, Katowice-Ligota, Poland) were kept...
in social cages with free access to fresh water and standard rodent food. It is important to emphasize that the experimental groups consisted of closely related animals (sisters) of the same age, which entered the hair cycle almost on the same day (in the case of depilation-induced cycling — exactly on the same day and hour).

We focused our attention on the first and the second postnatal hair cycle (Paus et al., 1999; Mueller-Roever et al., 2001), and the experimental groups were sacrificed on different time points after entering telogen. The time schedule of the experiments and symbols of groups are shown in Fig. 1. Only mice with homogenous telogen hair follicle population were selected for experiments (Fig. 2A, B). The telogen stage of the hair cycle was recognized by homogeneously pink color of the back skin, which can be appreciated particularly after depilation (Fig. 2B) (Paus et al., 1990; Slominski et al., 1991; 1994). All the experiments were approved by the Permanent Rector Committee for Bioethics of Experiments on Animals (Jagiellonian University permission number 303/97), and performed according to the international rules accepted by the Polish Parliament.

**Induction of hair growth.** We investigated telogen skin after spontaneously induced hair growth, but we also analyzed some animals in which telogen followed artificially induced anagen (Figs. 1B, E and 2B, C). The technique of anagen induction was based on the widely accepted depilation by application of melted 1:1 mixture of beeswax and gum rosin on the back skin of anesthetized animals (Ketamine, i.p. 0.75 mg/mouse, in PBS), and removal together with the hair shafts after hardening (Mueller-Roever et al., 2001).

**Hair, spleen and anagen skin preparation.** Mice in ketamine narcosis with whole back skin in telogen (Fig. 2A), were shaved and their hair shafts were kept for the further EPR analysis. Hair shafts of yellow gerbils (a 40 mg sample) served as a natural model material containing pheomelanin (Slominski et al., 2005; Plonka et al., 2003). The animals were killed by cervical dislocation, and spleens were removed. The presence, size and intensity of black spots (Fig. 2D–E) were estimated shortly before putting the organ into glass tubes, in which they were frozen, and after removal — measured by EPR. Finally, the mass of the spleens was recorded after the EPR measurement.

To obtain melanin-containing skin samples devoid of dead, highly pigmented and dry structures of hair shafts but consisting of both alive and melanized tissues, some mice in the late stage of depilation-induced anagen VI (day 15 after anagen in-

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**Figure 1.** Experimental design.

A–D. Telogen groups where spleen and hair shafts were examined. E. Group from which depilated anagen VI skin containing melanin was obtained. Empty circles — the day of anagen induction by depilation. Filled circles — the day of autopsy and sample harvesting. Vertical arrows — days when the last mice from the group entered telogen, as judged by homogenously pink color of the back skin (A–E), and the day of the onset of anagen VI (E).
duction, Fig. 1E) were anesthetized, depilated again (Fig. 2C), and sacrificed. The skin was tightly stuffed in standard glass tubes, frozen in liquid nitrogen, then pushed out from the tubes, and cut into two equal parts, so as to obtain samples of the same geometry (diameter 0.4 ± 0.02 cm; length 01.5 ± 0.1 cm) and mass (0.2 ± 0.003 g).

**Synthesis of dopa-melanin.** Dopa-melanin was prepared by autooxidation of D,L-dopa (Felix et al., 1978a). An aqueous solution of D,L-dopa was bubbled with air for 4 days at ambient temperature, with the pH kept constant at 8 (NH₄OH). Subsequently, the pigment formed was precipitated by lowering the pH to 3–3.5 (HCl), purified with several washings with re-distilled water, and dialysed against re-distilled, deionized water for 4 days. It is worth noticing here that the term “melanin solution” should rather be replaced with “melanin suspension”, because of difficulties in properly defining the type of solution formed by solubilized melanin (Crippa et al., 1989). The substance prepared by us contained about 10¹⁹ spins/mg dry mass, and it was used either in suspension (13 mg/ml, pH about 6.5) or as air-dry powder samples (8.2 ± 0.1 mg).

**EPR measurements and statistical analysis of the data.** All the samples were analyzed in a quartz Dewar filled with liquid nitrogen, at 77 K, using an EPR X-band (9.18 GHz) spectrometer Varian E3 with a TE 102 rectangular resonant cavity. Icicles of skin and dopa-melanin suspension were pushed out of the tubes directly into the Dewar. Powder samples of dopa-melanin were measured in glass capillaries sealed by melting and solidifying their both ends, and spleens were measured in a similar way to the skin samples. The skin and melanin suspension samples, but not spleen and the powder melanin samples, had the same geometry. Samples of shaved hair shafts (30 ± 0.5 mg) had been stuffed directly in the Dewar flask before liquid nitrogen was poured in. A powder sample of 1,1-diphenyl-2-picrylhydrazyl (DPPH) was used as a free-radical standard (g = 2.0037 ± 0.0002). All the measurements were performed at the magnetic field of 3260 ± 50 Gs. Other parameters of measurements were as follows: for qualitative comparison of the shape of the spectra (Fig. 3, Table 2) microwave power was 0.5 mW and modulation amplitude — 0.5 Gs. For qualitative comparisons (Table 1), modulation amplitude was raised to 5 Gs, and microwave power to 4 mW, whereas gain was changed to 250 000. To register the power saturation curves (Fig. 4, Table 2) the microwave power was changed every single dB over the range of 31–8 dB (0.05–12 mW), while the peak-to-peak amplitude of the signal was measured, and the
modulation amplitude was kept at 5 Gs. A lack of power saturation is usually manifested by a linear dependence of the signal amplitude on the square root of microwave power (Swartz & Swartz, 1983). Each spectrum was collected in a digital form (1024 points per spectrum) for 120 or 200 s at a constant of 0.1 or 0.3 s, 1–3 times (for the weak signals of spleens and yellow gerbil hair — 5–7 times) and averaged. To test the statistical significance of the observed differences, appropriate data were pooled, expressed as means ± S.E.M. or means ± S.D., and compared by two-tailed, independent Student’s t-test for populations of equal- and non-equal variations, which was tested using the F test.

RESULTS AND DISCUSSION

Splenic melanin predominates in early telogen

Murine C57BL/6 spleens (Fig. 2D) sometimes revealed black spots on their dorsal surface (Fig. 2E). No variability or gradation of the color intensity (e.g. from bright through dark-grey to black) could be observed, only variability in the size of the spots was found, which were always homogenously black. The presence of a black spot was not the rule, and it never corresponded absolutely to the type of hair cycle induction — in the studied groups of telogen mice after the depilation-induced or spontaneous hair growth (Table 1) there were always some spleens with the dark spots besides spleens completely devoid of the pigment. Even in the group of the highest percentage of melanotic spleens, there was one animal with the spleen completely devoid of the pigment, and vice versa — in groups revealing amelanotic spleens there were single animals with melanotic spleen. Nevertheless, spleens from mice which had just entered telogen (Group S1, S2, see Fig. 1A, C) revealed more frequently the presence of melanin (Table 1) than those from mice which had already been in telogen for several days (Group D2, S2(longer) — Fig. 1B, D). This observation represents the only correlation of the appearance of the dark spots with the hair cycling. This tendency was observed predominantly in animals with spontaneous hair cycling, which constituted 3 out of 4 experimental telogen groups (Fig. 1 A–D).

Although the most intensively pigmented spleen (of the highest amplitude of the free radical EPR signal, i.e. 5.675 a.u.) was found in the S2

| Number of the postnatal hair cycle and the way of its induction | Group designation | Duration of telogen until harvesting (day) | Total number of spleens | Number of spleens with a black spot (%) | Number of spleens without the black spots (%) | Free radical EPR signal amplitude; (a.u.), means from pigmented spleens ± S.D., or value for the only pigmented spleen in a group |
|----------------------------------------------------------------|------------------|------------------------------------------|-------------------------|------------------------------------------|---------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------|
| 1 spontaneous                                                  | S1               | 0                                        | 6                       | 3 (50%)                                  | 3 (50%)                                    | 3.08 ± 0.08                                                                                                                    |
| 2 spontaneous                                                  | S2               | 0                                        | 5                       | 4 (80%)                                  | 1 (20%)                                    | 3.52 ± 1.42                                                                                                                   |
| 2 depilation-induced                                           | D2               | 7                                        | 7                       | 1 (14.3%)                                | 6 (85.7%)                                  | 2.125                                                                                                                          |
| 2 spontaneous                                                  | S2(longer)       | 6                                        | 7                       | 1 (14.3%)                                | 6 (85.7%)                                  | 3.3                                                                                                                             |

Figure 3. Comparison of EPR spectra of synthetic and natural pigmented materials used in the experiments.

Note the singlet line in the region of the DPPH and dopa-melanin signal, present in the spectra of hair, skin and pigmented spleen of C57BL/6 mice, but not in the non-melanized organ, which confirms the melanin-like character of the black splenic pigment. The lack of hyperfine splitting (arrows) well appreciable in the spectrum of yellow gerbil hair indicates the eumelanin type of pigment contained in the spotted spleen. All spectra were registered at 77 K. Parameters of assay: see Material and Methods. Receiver gains: A — 40000, B — 300000, C — 1500000, D — 255000, E–F — 7500000, G — 750000.
group, which had 80% spleens with the spots (Table 1), it was hardly possible to correlate the melanin contents with the percentage of melanized spleens in particular groups. For example (Table 1), 3.3 a.u. was found for another spleen of group S2, for one spleen of group S1, and for the only pigmented spleen of group S2 (longer). The average mass of a pigmented spleen was slightly higher than the mass of non-melanized spleens, but this tendency was not significant ($P > 0.2$; not shown).

The splenic pigment may represent melanin produced in the spleen due to the same activators which initiate melanogenesis in skin, due to accumulation of circulating intermediates of melanogenesis and/or melanin collected and degraded in other sites of melanin synthesis. In the black C57BL/6 a/a mice the main source for such melanin is the pool of follicular melanocytes, which periodically undergo apoptosis in catagen (Tobin, 1998; Tobin et al., 1999). Yet the appearance of dark spots in spleens ought to be considered as a stochastic phenomenon. It is a “yes or no” phenomenon which cannot be inferred from any behavioral or morphological evidence observed in the mice. Without laparotomy only the probability that a particular mouse has pigmented spleen can be determined on the base of the time which passed after the onset of telogen.

**Spleen, hair shafts and anagen skin of C57BL/6 mice contain eumelanin**

Figure 3 shows EPR spectra of some pigmented materials obtained from rodents, including C57BL/6 mice, and the model melanins. In all the pigmented materials from mice, but not in non-pigmented spleens which revealed uniformly red coloration (Fig. 2D), a sharp singlet of a free radical character is seen in the region of both dopa-melanin and DPPH signals (Table 2, Fig. 3). Also the power saturability of the melanin signals is strong (Table 2, Fig. 4), which indicates their free-radical character (Felix et al., 1978b; Sarna & Hyde, 1978).

The signal appeared as a slightly asymmetric singlet placed around $g = 2.003–2.004$, of the peak-to-peak line-width of about 4.5–5.5 Gs, without any hyperfine splitting, which might indicate contribution of pheomelanin (Sealy et al., 1982; Slominski et al., 2005). Therefore, we can conclude that the type

![Figure 4. Saturation of the melanin EPR signal with microwave power.](image)

A–B. Dopa melanin. C–D. Biological materials from C57BL/6 mice comprising normally hydrated tissues (C, depilated anagen VI skin; D, spleen with melanotic spot). E. Dry black hair shafts from C57BL/6 mice. All measurements were carried out at 77 K. Parameters of assay: see Materials and Methods. Receiver gains: A–B — 2000, C — 125000, D — 400000, E — 15000. F. Superposition of curves A–E normalized for the same value of amplitude in maximum to compare the shapes of the saturation curves.
Paramagnetic properties of splenic eumelanin indicate its heterogeneity and partial degradation

Table 2. Parameters of the EPR signal of samples of pigments and biological materials obtained from C57BL/6 mice: peak-to-peak linewidth (ΔH) and the microwave saturation power at which the saturation curve reaches its maximum (means of 3–6 samples ± S.E.M.).

| Material                  | ΔH [Gs]     | Maximum of the power saturation curve [mW] (dB) |
|---------------------------|-------------|-----------------------------------------------|
| Dopa melanin (powder)     | 4.74±0.11   | 0.61±0.04                                      |
| Dopa melanin (suspension) | 4.45±0.12   | 1.01±0.27                                      |
| Depilated anagen VI skin  | 5.52±0.05   | 0.27±0.01                                      |
| Hair shafts               | 5.56±0.07   | 0.29±0.01                                      |
| Spleens with black spots  | 5.00±0.07   | 0.84±0.46                                      |

The difference between ΔH for spleens and other murine materials was highly significant (0.01 > P ≥ 0.001), and between the position of maxima — significant (0.05 > P ≥ 0.01). Parameters of assay performed at 77 K: see Fig. 3, 4, and Material and Methods.

Of melanin contained in the black-spotted spleens is eumelanin, the same as in the hair shafts of the animals. To our knowledge this is the first EPR confirmation of the presence of eumelanin in spleens of C57BL/6 mice.

Power saturability is also different — the saturation curves were in general much flatter in the case of spleens than for hair shafts or skin (Fig. 4C, D, E). The placement of the maximum was also shifted towards higher power, as compared to the hair/skin melanins. The poorer saturability and the flatter saturation curves of pigmented spleens, as compared to pigmented skin and hair shafts suggest higher heterogeneity of the local environments of the splenic paramagnetic centers (Swartz & Swartz, 1983), and closer interactions with some rapidly relaxing paramagnetic centres, like dioxygen, iron (II), iron (III) or copper (II), which betrays heterogeneity within the pool of splenic melanin itself (Felix et al., 1978a). This finding supports the view that at least a fraction of splenic melanin undergoes degradation. It must be emphasized that, as the paramagnetism of splenic melanin is preserved, this degradation may primarily concern the proteins which are present along with melanin in melanosomes (Okazaki et al., 1985), and which are probably degraded first (Borovansky & Elleder, 2003; Sulaimon & Kitchell, 2003). Proteins may influence the linewidth as well as the power saturability (Sarna & Hyde, 1978; Okazaki et al., 1985). This may to some degree explain the narrower signals of splenic melanin, which was also the case for the protein-free dopa-melanin, as compared to natural melanins. Proteolysis can be followed by oxidative degradation of melanin itself (Stanka et al., 1988; Borovanský & Elleder, 2003).

Among the putative candidates for the function of “melanophages” responsible for transport and/or degradation of skin/hair follicle melanin, Langerhans cells seem the most important ones, due to their considerable migration potential and the role in scavenging the apoptotic bodies in catagen (Tobin, 1998; 1999). Other important cells potentially responsible for this phenomenon may be macrophages, due to their presence in hair follicles and their role in catagen remodeling (Parakkal, 1969a; 1969b). Their importance may increase with the aging of the animal (van der Heijen et al., 1995). Another type of cells suspected of transiently depositing ectopic melanin and particularly active during premature catagen induced by chemotherapy are noncortical keratinocytes (Tobin et al., 1999). Finally, macrophages and leukocytes (neutrophils and lymphocytes), have been suggested to participate in the process of “alternative melanin transport” in the vertebrate organism (Wassermann, 1967) to the place of its visceral deposition.

To our surprise, there were no evident qualitative differences between melanin from depilated anagen VI skin (Fig. 4C, Table 2) and that from hair shafts (Fig. 4E, Table 2), although we expected that the state of dehydration of hair melanin, and the possible presence of oxygen, as compared with the melanosomes in live melanocytes and keratinocytes,
might to some degree influence the spin-lattice relaxation properties of melanin paramagnetic centers, which might be reflected in the power saturability of the signals, and their linewidths (Sarna & Hyde 1978, Felix et al., 1978b; Swartz & Swartz 1983). This finding may be important for proper monitoring of the on-going melanogenesis in anagen of depilation-induced murine skin by EPR, in particular when the dry hair shaft is being produced (anagen IV–V) and stretches out of the skin (anagen VI) (Slominski et al., 1994; 1996; Plonka et al., 1995).

CONCLUSIONS

Using EPR spectroscopy we demonstrated here the presence of eumelanin in black-spotted spleens of C57BL/6 mice with similar biophysical properties as the pigment of black hair shafts and anagen skin melanocytes. However, the exact nature of the splenic eumelanin is different from that of skin/hair shaft melanin and reveals more heterogeneity, which can be to some degree explained by pigment degradation. The existence of melanin in the spleen is not the rule, although spleens of mice which had just entered telogen tended to contain pigmented spleens more often than was the case in animals which had been in telogen for a longer time.

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