An anaerobic mitochondrion that produces hydrogen

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subject to the constraints that no lichen covered the trunk and no young trees with a trunk circumference less than 0.9 m were used. Colour matches of treatments to natural bark were verified by spectrophotometry of stimuli and bark, followed by modelling of predicted photon catches of a typical passerine bird, the blue tit’s (Parus caeruleus) single cone photoreceptors, with irradiance spectra from overcast skies in the study site. Our acceptance criterion was simply that cone captures for the experimental stimuli fell within the measured range of those for oak bark.

Experiment 1 used black patterns printed onto dark brown card. Patterns were samples of digital photos of the oak trees at 1:1 reproduction, converted using ImageJ to grey scale and thresholded at 50% to binary (black/white) images to provide, when printed onto different trees, were used for each replicate target.

Experiment 2 used bicoloured targets printed onto waterproof paper (Hewlett Packard Laserjet Tough Paper) with a Hewlett Packard Colour Laserjet 2500 (600 dots per inch) printer, with colour pairs chosen to have either high or low contrast. Colours were chosen from frequency distributions of the eight-bit RGB (red, green, blue) values from digital photographs of the oak trees in the study site, reduced to 16 bins in each colour channel. Photos (about 267 mm × 200 mm; 2,560 pixels × 1,920 pixels) were taken with a Nikon Coolpix 5700 camera, calibrated to linearize the relationship between radiance and the grey scale in each colour channel, and saved as uncompressed TIFF files. Digital photographs lack ultraviolet information that birds can see, but lichen-free oak bark reflects negligible ultraviolet. Even a properly calibrated RGB image does not precisely simulate the avian-perceived colour of many natural objects, owing to differences in the spectral sensitivity of bird long-wave, medium-wave and short-wave cones compared with human cones. However, because our treatments varied only in relative colour contrast, any error arising from this method was considered minor, an assumption verified retrospectively by spectrophotometry and colour-space modelling. We chose colour pairs from the eight most frequent RGB triplets in the bark photos as follows: a ‘background’ colour, then a triplet that was similar to the background (low contrast), and one that differed markedly (high contrast). The major difference between colours was in overall brightness not hue, but we could not systematically vary only one colour dimension within the available common bark colours. Sample numbers of background and contrasting colours were balanced for which was darker/lighter, and so there were no significant differences between bicoloured treatments in the brightest or darkest colour or average colour (analyses of variance on RGB sums and all possible ratios; P > 0.9). Monochrome treatments were also created as the means of the respective R, G and B values of the two colours in bicoloured high-contrast and low-contrast treatments. Different colour pairs and patterns, from different trees, were used for each replicate target.

Survival analysis was by Cox regression with the factors treatment and block. Cox regression assumes that all survival functions have the same shape; this proportional hazards assumption was checked by plotting partial residuals against ranked survival times. There were significant block effects in both experiments (in experiment 1, the hazards assumption was checked by plotting partial residuals against ranked survival times24. There were significant block effects in both experiments (in experiment 1, χ² = 141.78, d.f. = 9, P < 0.001; in experiment 2, Wald = 271.50, d.f. = 9, P < 0.001), reflecting differences in average predation rates in different parts of the woods on different dates, but this was not relevant to our hypotheses.

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Authors’ contributions I.C.C. designed the experiments and stimuli; M.S., J.S., T.M. and I.C.C. performed the experiments; A.P. wrote the programs for colour analysis and camera calibration; T.S.T. advised on design and colour modelling.

Competing interests statement The authors declare that they have no competing financial interests.

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An anaerobic mitochondrion that produces hydrogen

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Hydrogenosomes are organelles that produce ATP and hydrogen, and are found in various unrelated eukaryotes, such as anaerobic flagellates, chytridiomycete fungi and ciliates. Although all of these organelles generate hydrogen, the hydrogenosomes from these organisms are structurally and metabolically quite different, just like mitochondria where large differences also exist. These differences have led to a continuing debate about the evolutionary origin of hydrogenosomes. Here we show that the hydrogenosomes of the anaerobic ciliate Nyctotherus ovalis, which thrives in the hindgut of cockroaches, have retained a rudimentary genomic encoding components of a mitochondrial electron transport chain. Phylogenetic analyses reveal that these proteins cluster with their homologues from aerobic ciliates. In addition, several nucleus-encoded components of the mitochondrial proteome, such as pyruvate dehydrogenase and complex II, were identified. The N. ovalis hydrogenosome is sensitive to inhibitors of mitochondrial complex I and produces succinate as a major metabolic end product—biochemical traits typical of anaerobic mitochondria. The production of hydrogen, together with the presence of a genome encoding respiratory chain components, and biochemical
Figure 1 A 14,027-bp fragment (mtg 1) of the hydrogenosomal genome of *N. ovalis* var. *Blaberus* Amsterdam. Black boxes, RNA coding genes; shaded boxes, genes with significant similarity to mitochondrial genes; white boxes, unknown ORFs (named according to the number of codons); arrows, cDNAs identified so far. The numbers indicate the nucleotide positions on the 14-kb clone (mtg 1). The longest ORF (4,179–9,728) contains a stretch with significant similarity to *nad5*. A potential start codon for a putative *nad5* transcript is marked with an asterisk.

Figure 2 Phylogenetic analysis of hydrogenosomal genes. Both the organellar 12S (SSU) rRNA gene (b) and the nuclear hsp60 (c) reveal a ciliate ancestry for the hydrogenosome of *N. ovalis*. The same is true for the components of a "mitochondrial" complex I, the *nad7* (49 kDa; organellar, a) and 51 kDa (nuclear, d) genes. The phylogenies were derived using MrBayes and neighbour joining: the topologies correspond to the maximum-likelihood (MrBayes) approach, and the values at the nodes indicate the posterior probability for the partition and its bootstrap value, respectively. Only values higher than 50% are indicated. See Supplementary Information. EB, Eubacteria.


features characteristic of anaerobic mitochondria, identify the *N. ovalis* organelle as a missing link between mitochondria and hydrogenosomes.

Hydrogenosomes and their highly reduced relatives, mitosomes, generally lack an organelle genome\(^8\)-\(^9\), hampering clarification of their origin. Two models for the origin of hydrogenosomes are currently debated. The first postulates that the ancestral mitochondrial endosymbiont gave rise to aerobically functioning mitochondria, which subsequently evolved into hydrogenosomes by the acquisition of genes encoding enzymes essential for an anaerobic metabolism\(^1\)^-\(^3\). The second hypothesis presumes that hydrogenosomes and mitochondria originated from one and the same ancestral—facultatively anaerobic—(endo)symbiont, followed by specialization to aerobic and anaerobic niches during eukaryotic evolution\(^1\)^-\(^4\). To address this issue we investigated DNA in hydrogenosomes of *N. ovalis*, which was previously identified by immunocytochemical methods\(^5\). Intact *N. ovalis* hydrogenosomes isolated by cell fractionation contained DNA between 20 and 40 kilobases (kb) long. Long-range polymerase chain reaction (PCR) with this DNA with the use of specific primers for the hydrogenosomal small-subunit (SSU) ribosomal RNA\(^1\) and *nad7* (obtained earlier by PCR with degenerated primers) yielded a 12-kb fragment of the organellar genome. It encodes four genes of a mitochondrial complex I (*nad2, nad4L, nad5* and *nad7*), two genes encoding mitochondrial ribosomal proteins RPL 2 and RPL 14, and a \(^1\)RNA gene (Fig. 1). *nad2* and *nad4L*, which are generally poorly conserved among ciliates, could be identified by using multiple sequence alignments and an analysis of their membrane-spanning domains as described.

### Table 1 Nyctotherus ovalis genes encoding mitochondrial proteins and RNAs

<table>
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<tr>
<th>Type</th>
<th>Gene product</th>
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<th>Codon use</th>
<th>cDNA Target</th>
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<td>RPL 20</td>
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<td>Nuc</td>
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<td>N</td>
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<td>Nuc</td>
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<td></td>
<td>Heat shock protein HSP92 YMR186W_Chr</td>
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<td>Nuc</td>
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<td>No</td>
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<td>Nuc</td>
<td>? AJ871330</td>
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<td>(Protein YDJ1)</td>
<td>N</td>
<td>Nuc</td>
<td>? AJ871331</td>
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</table>

For a complete table see Supplementary Information. N, nucleus; H, hydrogenosome; nuc, nuclear; mt, mitochondrial; mtg 1, 12-kb clone of hydrogenosomal genome (AJ871267); ?, low-probability support so far because full-length cDNAs have not yet been isolated (the N terminus might be incomplete, or it might contain an in-frame intron or alternative start codons). Accession numbers for cDNAs: * AJ871574 and AJ871575; † AJ871576; ‡ Y608633 and Y608634; § Y619800 and Y619801; ¶ Y623917; ↔ Y6239023; †† Y6068627 and Y6068634.

### Table 2 Glucose metabolism of Nyctotherus ovalis

<table>
<thead>
<tr>
<th>Labeled end products</th>
<th>([\text{U}-^{14}\text{C}]\text{glucose})</th>
<th>([\text{[6}-^{14}\text{C}]\text{glucose})</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>((\mu\text{mol h}^{-1}) per mg protein)</td>
<td>((\mu\text{mol h}^{-1}) per mg protein)</td>
</tr>
<tr>
<td>Acetate</td>
<td>427</td>
<td>467 ± 87</td>
</tr>
<tr>
<td>Lactate</td>
<td>220</td>
<td>156 ± 116</td>
</tr>
<tr>
<td>Succinate</td>
<td>112</td>
<td>79 ± 65</td>
</tr>
<tr>
<td>Ethanol</td>
<td>44</td>
<td>29 ± 26</td>
</tr>
<tr>
<td>O₂</td>
<td>204</td>
<td>ND</td>
</tr>
<tr>
<td>Formate</td>
<td>ND</td>
<td>ND</td>
</tr>
</tbody>
</table>

Cells were incubated for 48 h at 25°C in micro-aerobic conditions in medium with either \([\text{U}-^{14}\text{C}]\text{glucose}\) or \([\text{[6}-^{14}\text{C}]\text{glucose}\). Excreted end products are shown as means ± s.d. of three independent experiments \((\text{[6}-^{14}\text{C}]\text{glucose})\) or as the means of two independent experiments \((\text{[U}-^{14}\text{C}]\text{glucose})\). Other excreted end products were less than 2% of the total excreted end products. ND, not detectable.

*Percentage of the total of acetate, lactate, succinate and ethanol.

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Phylogenetic analysis revealed clustering of these genes with their homologues from the mitochondrial genomes of aerobic ciliates (Fig. 2, and Supplementary Information). All genes exhibit a characteristic mitochondrial codon-usage and lack amino-terminal extensions that could function as a mitochondrial targeting signal (Table 1). Complementary DNAs isolated for \textit{nad5} and \textit{nad7} show that they are transcribed. Translation with a nuclear genetic code from \textit{N. ovalis}, rather than the ciliate mitochondrial code, leads to numerous stop codons (not shown). Five additional open reading frames (ORFs 236, 262, 71, 161 and 199) do not show significant sequence similarity to ORFs from the mitochondrial genomes accessible in the EMBL database. Two ORFs overlap with neighbouring ORFs as in other mitochondrial genomes\textsuperscript{12}.

Macronuclear gene-sized chromosomes encoding the 24-kDa, 51-kDa and 75-kDa subunits of mitochondrial complex I and the Fp and Ip subunits of mitochondrial complex II were cloned with a PCR-based approach. These have a nuclear codon usage, are transcribed (Table 1), encode a putative N-terminal mitochondrial targeting signal and branch with their mitochondrial homologues from aerobic ciliates in phylogenetic analyses (Fig. 2, Table 1 and Supplementary Information). They are similar to the two complex I-like Ndh51 and Ndh24 proteins discovered in \textit{Trichomonas vaginalis}\textsuperscript{18,19}, because a phylogenetic analysis including the mitochondrial homologues from \textit{N. ovalis} and certain aerobic ciliates reveals that all these proteins belong to a cluster of mitochondrial complex I homologues (see Supplementary Information). Thus, in \textit{N. ovalis}, 7 of the 14 genes encoding core proteins of mitochondrial complex I, and two of the four proteins of mitochondrial complex II, have been identified so far. They are well conserved, are transcribed, and cluster with the mitochondrial homologues of their aerobic (ciliate) relatives, indicating that the hydrogenosomes of \textit{N. ovalis} have retained parts of a functional mitochondrial electron-transport chain.

Hydrogenosomes of \textit{N. ovalis} have typical mitochondrial cristae and contain cardiolipin\textsuperscript{11}. They are closely associated with endosymbiotic methanogens, which are biomarkers for hydrogen formation by the \textit{N. ovalis} hydrogenosomes\textsuperscript{20}(Fig. 3a). The organelles stain with Mitotracker Green FM and fluoresce with rhodamine 123, indicating the presence of a membrane potential (Fig. 3). Carbonyl cyanide \textit{p}-trifluoromethoxyphenylhydrazone (FCCP) (5 \textmu M) prevented staining with rhodamine 123, indicating the possible presence of a proton gradient. Moreover, staining of the hydrogenosomes with rhodamine 123 was also prevented after incubation of the ciliates with rotenone, piericidin, fenazaquin and 1-methyl-4-phenylpyridinium (MPP\textsuperscript{+}) (classical inhibitors of mitochondrial complex I (ref. 21)), but not with cyanide (1 mM) or antimycin A (inhibitors of mitochondrial complex III and IV; Fig. 3). Similarly, treatment with cyanide and salicylhydroxamic acid stains the same organelles. The inserts in \textbf{d} and \textbf{e} (outside the ciliate) show the organelles seen in the box inside the ciliate, \textbf{f}, \textbf{h}, \textbf{i}, incubation of living cells with inhibitors of mitochondrial complex I (MPP\textsuperscript{+}, \textbf{f}, piericidin \textbf{h} and rotenone \textbf{i})\textsuperscript{21} completely prevents staining of the organelles by R123. \textbf{g}, incubation with cyanide (1 mM) or antimycin A (not shown) does not interfere with staining by R123. For additional information see the text and Supplementary Information. Scale bars, 1 \mu m (\textbf{a}, also applies to \textbf{b}); 10 \mu m (\textbf{c}–\textbf{i}).

Figure 3 Hydrogenosomes of \textit{N. ovalis} exhibit complex I activity. \textbf{a}, \textbf{b}, Electron micrographs of a hydrogenosome of \textit{N. ovalis} (\textbf{a}) and a mitochondrion of \textit{Euplotes} sp. (\textbf{b}). White arrowheads mark cristae; \textit{m}, endosymbiotic methanogenic archaeon\textsuperscript{15,20}. \textbf{c}, Fluorescence picture of \textit{N. ovalis} hydrogenosomes (bright dots), which were released from the cell by gentle squashing after being stained \textit{in vivo} with ethidium bromide. \textbf{d}, Rhodamine 123 (R123) also stains the hydrogenosomes, the only organelles matching the expected size (compare \textbf{a}, and Supplementary Information). \textbf{e}, Mitotracker green FM stains the same organelles. The inserts in \textbf{d} and \textbf{e} (outside the ciliate) show the organelles seen in the box inside the ciliate. \textbf{f}, \textbf{h}, \textbf{i}, Incubation of living cells with inhibitors of mitochondrial complex I (MPP\textsuperscript{+}, \textbf{f}, piericidin \textbf{h} and rotenone \textbf{i})\textsuperscript{21} completely prevents staining of the organelles by R123. \textbf{g}, Incubation with cyanide (1 mM) or antimycin A (not shown) does not interfere with staining by R123. For additional information see the text and Supplementary Information. Scale bars, 1 \mu m (\textbf{a}, also applies to \textbf{b}); 10 \mu m (\textbf{c}–\textbf{i}).
acid (SHAM), inhibitors of mitochondrial complex IV of the respiratory chain and the plant-like alternative oxidase known from certain mitochondria3, respectively, neither killed *N. ovalis* nor interfered with its oxygen consumption under aerobic conditions (not shown). These observations not only indicate the absence of a functional complex III and IV and the absence of a terminal (plant-like) alternative oxidase, but also reveal the presence of a functional mitochondrial complex I as the source of the organellar proton gradient4. The oxygen consumption of *N. ovalis* observed under aerobic conditions is most probably a detoxification mechanism, and longer exposure to atmospheric oxygen kills the ciliates effectively.

Metabolic experiments using tracer amounts of uniformly labelled (U-)14C-glucose revealed that *N. ovalis* catabolizes glucose predominantly into acetate, lactate, succinate and smaller amounts of ethanol, in addition to CO2 (Table 2). The presence of oxygen did not cause significant changes in the pattern of excreted end products (not shown). Notably, incubations in the presence of [6-14C]glucose did not result in the formation of labelled CO2. Because 14C-labelled CO2 is released from [6-14C]glucose by successive decarboxylations through multiple rounds in the Krebs cycle, the absence of labelled CO2 after application of [6-14C]glucose indicates the absence of a complete Krebs cycle. The observed excretion of 14C-labelled CO2 after incubation with [U-14C]glucose could be the result of either pyruvate dehydrogenase (PDH) activity, as in typical aerobic mitochondria, or pyruvate:ferredoxin oxidoreductase (PFO) activity, as in the hydrogenosomes of *T. vaginalis*. A third possibility for pyruvate catabolism, pyruvate formate lyase activity22,23, can be excluded because no detectable amounts of formate were produced from [U-14C]glucose (Table 2). We failed to identify genes for PFO but succeeded in isolating three of the four PDH genes, namely the E1α, E1β and E2 subunits, which are expressed as cDNA, indicating that *N. ovalis* uses a mitochondrial PDH for oxidative decarboxylation. Significant amounts of 14C-labelled succinate from both [U-14C]glucose and [6-14C]glucose (Table 2) indicate that endogenously produced fumarate is used as a terminal electron acceptor, as in some anaerobic mitochondria3. Fumarate reduction in *N. ovalis* (to account for the production of succinate) is most probably catalysed by a membrane-bound complex II (see above; Table 1, and Supplementary Information), which is coupled to complex I through electron transport mediated by quinones. Mass spectrometry coupled to liquid chromatography of lipid extracts from *N. ovalis* revealed the presence of small amounts of quinones (rhodiquinone 9 and menaquinone 8) at a concentration of about 1 pmol per mg protein (Supplementary Information). This amount is at least two orders of magnitude lower than in other eukaryotes known to possess anaerobic mitochondria producing succinate4. The low concentration of quinones in *N. ovalis* cells might reflect the intermediate state of their hydrogenosomes, occupying a position between mitochondria (which contain a membrane-bound electron transport chain) and previously characterized hydrogenosomes (which do not)1,4,13,14.

Although an F1-F2-ATP synthase has not yet been identified, the hydrogenosome of *N. ovalis* has retained certain basal energy-generating functions of an aerobically functioning mitochondrion1. To explore the presence of additional ‘mitochondrial’ traits in *N. ovalis*, we performed a reciprocal Smith–Waterman sequence comparison between about 2,000 six-frame-translated clones from our genomic DNA library of *N. ovalis* and the yeast15 and human16 mitochondrial proteins. We identified 53 additional nuclear genes encoding potential mitochondrial proteins in addition to components of the mitochondrial import machinery (Table 1, and Supplementary Information).

In contrast, the hydrogenase of *N. ovalis* does not exhibit any mitochondrial traits. This hydrogenase is rather unusual in comparison with other eukaryotic hydrogenases because it seems to be a fusion of a [Fe] hydrogenase with two accessory subunits of different evolutionary origin1,15. These subunits should allow NADH reoxidation in combination with the [Fe] hydrogenase, because they exhibit a significant sequence similarity to the *hox F* and *hox U* subunits of β-proteobacterial [Ni–Fe] hydrogenases, in contrast to similar, recently described hydrogenosomoprotal proteins (24 and 51 kDa) of putative mitochondrial origin from *T. vaginalis*16,19. The ‘mitochondrial’ 24-kDa and 51-kDa genes of *N. ovalis* are clearly different from the above-mentioned hydrogenase modules and are likely to function in mitochondrial complex I (Fig. 3, Table 1, and Supplementary Information). Moreover, the catalytic centre of the hydrogenase (the H-cluster) clusters neither with any of the hydrogenases of *Trichomonas* sp., *Furmocyes* sp. and *Neocallimastix* studied so far, nor with any of the hydrogenase-related Nar proteins, which seem to be shared by all eukaryotes. Rather, the *N. ovalis* hydrogenase is more closely related to [Fe] hydrogenases from β-proteobacteria1,13,15. These observations suggest that the hydrogenase of *N. ovalis* has been acquired by lateral gene transfer.

It should be realized that the hydrogenosome of *N. ovalis* is so far unique and not representative of all hydrogenosomes, which seem to have been involved repeatedly and independently—albeit from the same ancestral mitochondrial-type organelle. All our data identify the hydrogenosome of *N. ovalis* as a ciliate-type mitochondrion that produces hydrogen. The presence of respiratory-chain activity of mitochondrial complex I and II, in combination with hydrogen formation, characterizes the *N. ovalis* hydrogenosome as a true missing link in the evolution of mitochondria and hydrogenosomes.
Smart Race cDNA amplification kit (Clonetech). Expressed sequence tags were amplified by PCR with the universal adapter primer provided with the kit and the various, specific internal primers.

Complete macronuclear gene-sized chromosomes

Telomere-specific primers in combination with internal gene sequences allow a straightforward recovery of the complete gene. The specific [internal] primers were based on the DNA sequences of internal fragments of the various genes, which were recovered previously by PCR with degenerated primers for conserved parts of the various genes.

Phylogenetic analysis

Protein sequences were aligned with ClustalW and Muscle; unequivocally aligned positions were selected with Gblocks. HMMs were inferred with maximum likelihood by using a discrete gamma-distribution model with four rate categories plus invariant positions and the Poisson amino acid similarity matrix, and maximum likelihood by using a discrete gamma-distribution model with four rate categories plus invariant positions and the Poisson amino acid similarity matrix, and neighbour joining as implemented in ClustalW, correcting for multiple substitutions with the Gonnet amino acids identity matrix, and bootstrapping with 100 samples. ORFs with a lower size limit of 100 nucleotides were identified with ORF Finder (http://www.ncbi.nlm.nih.gov/entrez/query.fcgi). tRNAs were identified with tRNAscan-SE (http://www.genetics.wustl.edu/edin/tRNAscan-SE). Potential mitochondrial import signals were detected with MITOP (http://mips.gsf.de/cgi-bin/proj/medgen/mitofinder).

Supplementary Information

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Competing interests statement

The authors declare that they have no competing financial interests.

Correspondence and requests for materials should be addressed to J.H.P.H. (j.hackstein@science.ru.nl). Sequences have been submitted to the EMBL database under accession numbers AFA80921, AJ871267, AJ871334, AJ871261, AJ871573–AJ871576, AY608627, AY608632–AY608634, AY616150–AY616152, AY619980, AY623917, AY623919, AY623925, AY629296, AY626883, AY626884, AY626888.

The perception of surface albedo (lightness) is one of the most basic aspects of visual awareness. It is well known that the apparent lightness of a target depends on the context in which it is embedded1–4, but there is extensive debate about the computations and representations underlying perceived lightness. One view asserts that the visual system explicitly separates surface reflectance from the prevailing illumination and atmospheric conditions in which it is embedded5–10, generating layered image representations. Some recent theory has challenged this view and asserted that the human visual system derives surface lightness without explicitly segmenting images into multiple layers11,12. Here we present new lightness illusions—the largest reported to date—that unequivocally demonstrate the effect that layered image representations can induce dramatic lightness illusions, causing identical texture patches to appear either black or white. These results indicate that mechanisms involved in decomposing images into layered representations can play a decisive role in the perception of surface lightness.

The amount of light projected to the eyes (luminance) is determined by a number of factors: the illumination that strikes visible surfaces, the proportion of light reflected from the surface and the amount of light absorbed, reflected or deflected by the prevailing atmospheric conditions (such as haze or other partially transparent media). Only one of these factors, the proportion of light reflected (lightness), is associated with an intrinsic property of