Binocular pattern deprivation interferes with the expression of proteins involved in primary visual cortex maturation in the cat

Karolina Laskowska-Macios1,2, Julie Nys2, Tjing-Tjing Hu2, Monika Zapasnik1, Anke Van der Perren3, Malgorzata Kossut1, Kalina Burnat1 and Lutgarde Arckens2*

Abstract

Background: Binocular pattern deprivation from eye opening (early BD) delays the maturation of the primary visual cortex. This delay is more pronounced for the peripheral than the central visual field representation within area 17, particularly between the age of 2 and 4 months [Laskowska-Macios, Cereb Cortex, 2014].

Results: In this study, we probed for related dynamic changes in the cortical proteome. We introduced age, cortical region and BD as principal variables in a 2-D DIGE screen of area 17. In this way we explored the potential of BD-related protein expression changes between central and peripheral area 17 of 2- and 4-month-old BD (2BD, 4BD) kittens as a valid parameter towards the identification of brain maturation-related molecular processes. Consistent with the maturation delay, distinct developmental protein expression changes observed for normal kittens were postponed by BD, especially in the peripheral region. These BD-induced proteomic changes suggest a negative regulation of neurite outgrowth, synaptic transmission and clathrin-mediated endocytosis, thereby implicating these processes in normal experience-induced visual cortex maturation. Verification of the expression of proteins from each of the biological processes via Western analysis disclosed that some of the transient proteomic changes correlate to the distinct behavioral outcome in adult life, depending on timing and duration of the BD period [Neuroscience 2013;255:99-109].

Conclusions: Taken together, the plasticity potential to recover from BD, in relation to ensuring restoration of normal visual input, appears to rely on specific protein expression changes and cellular processes induced by the loss of pattern vision in early life.

Background

Previous investigations emphasizing the molecular development of primary visual cortex typically dealt with its central visual field representation, or did not take a distinction between the central and peripheral visual field representations into account. Age-dependent expression profiles of proteins involved in neurite outgrowth, energy metabolism, synaptic development and neurotransmission are described [1–7]. Yet, when considering ocular dominance plasticity or synapse formation the development of the peripheral visual field representation in cat area 17 is slower than that of the central visual field representation [8, 9–12] (reviewed in [11]). In area 17 of marmoset monkey, Bourne and colleagues [13] showed that neurofilament protein patterns correlate with the maturation state of a given neocortical brain region and that the peripheral visual field representation in area 17 also achieves such a mature pattern later than the central visual field representation does.

In cat, lack of patterned visual input has been shown to differentially affect the development of the X- and Y-dominated functional pathways carrying information predominantly derived from the central versus the peripheral visual field respectively. At the level of the retina, early binocular pattern deprivation (BD), an animal model of congenital cataract, if applied for 6 months, results in permanent changes in the number and dendritic tree stratification of Y-type motion-sensitive alpha retinal
ganglion cells [14]. At the level of the thalamus, in the LGN, BD specifically affects the development of the Y-cell pathway, processing motion-sensitive visual information, but not the X-cell pathway, dominant in processing high resolution visual information (for review see [15]). Importantly, the parallel motion perception impairments observed in cat have also been described in patients with congenital binocular cataract (cat: [16, 17, 18]; human: [19, 20]). Furthermore, at the level of the visual cortex, early BD delays the maturation of peripheral area 17 to a larger extent than its central counterpart, as visualized by age- and cortical region-dependent expression changes for the activity reporter gene zf268 in the visual cortex of BD kittens [8].

We therefore decided to apply a functional proteomics approach to find molecular correlates for the centro-peripheral developmental gradient in area 17 in order to identify important proteins underlying cortical-region specific maturation. To this end, we separately assessed the central and peripheral region of area 17 of 2- and 4-month old BD (2BD and 4BD; early onset BD) kittens and age-matched controls with normal visual experience. Age and the distinct BD-induced delay of cortical maturation were thus considered as factors influencing protein expression in relation to cortical maturation. Two-Dimensional Difference Gel Electrophoresis (2-D DIGE) combined with mass spectrometry and Ingenuity Pathway Analysis (IPA) allowed the prediction of relevant molecular pathways and biological processes. To validate these proteomics observations and to investigate and compare protein expression profiles in additional experimental conditions, Western analysis was also applied to homogenates from area 17 of 6-month old BD kittens, a late onset BD group binocularly deprived during the 3rd and 4th month of age after 2 months of normal vision (2N 2BD) and extra normally sighted controls of 1 and 6 months, and 2 years (Adult). Analysis of the developmental profiles of protein expression in the context of normal visual stimulation, early onset as well as late onset BD enabled us to determine if BD regulation of protein expression depended on time of onset of BD or not.

We demonstrate how specific developmental protein expression changes are postponed especially in the peripheral visual field representation under BD. In particular, early BD exerts an influence on protein expression in a direction suggestive of a negative regulation of neurite outgrowth, synaptic transmission and clathrin-mediated endocytosis.

**Results**

**2-D DIGE screening for protein expression changes related to cortical maturation**

Age, BD, and centro-peripheral expression differences were considered valid parameters to chart molecular events in relation to cortical maturation (Fig. 1a-c, Table 1).

**Analysis at spot level**

Analysis of all the spot expression patterns revealed a total of 39 differentially expressed spots (Fig. 1b). When visualizing the effect of age on the number of differential spots for central and peripheral area 17 for normal and BD animals in a Venn diagram (Fig. 1c), a first indication for a reduced effect of age on peripheral area 17 becomes apparent with only 4 differential spots for peripheral area 17 of BD cats versus 7 for normal subjects.

**Analysis at protein level**

Table 1 summarizes the results of a similar analysis upon mass spectrometric identification of the 36 proteins in 32 spots (see Material and Methods, Additional file 1: Table S1).

**Normal**

Comparison of the protein expression patterns within the central or the peripheral regions in area 17 between kittens of 2 and 4 months revealed an age-dependent expression level for 20 proteins (grey box). Only four proteins (rows 3, 4, 8 and 19) were similarly regulated in the central and peripheral region, whereas seven proteins were specific for central and nine for peripheral area 17.

**BD**

When probing for the effect of early BD on these age-specific expression patterns, only for CRMP2 and CRMP4 the expression changed in the same direction for both regions (rows 2 and 3). Eighteen out of twenty proteins did not show the same age-dependent modulation as in normal kittens. Instead, the factor age identified five additional proteins in BD kittens, not differential between 2N and 4N (rows 21–25). Comparison of the BD-subjects with their age-matched normal controls (columns 2BD/2N and 4BD/4N) revealed an extra set of 11 proteins (rows 26–36) with an expression deviating from normal levels, resulting in a list of 36 differential proteins with a potential role in cortical maturation.

In general, age-dependent protein expression changes occurred less frequently in peripheral than in central area 17 under BD (five versus nine proteins; column BD/age) indicating that protein expression changes relevant to area 17 maturation may be postponed, especially in the peripheral region. Likewise, centro-peripheral differences in protein expression were observed only at 2 months in normal animals (proteins in rows 9, 12, 13 and 18 were upregulated in the peripheral region; 19 and 36 upregulated in the central region; Additional file 1: Table S1), whereas in BD animals proteins showed a centro-peripheral expression gradient for both age groups (three proteins in 2BD and four in 4BD; indicated by ‘+’ in columns 2BD/2N and 4BD/4N). Importantly, when comparing expression of all identified proteins between 4BD and younger normal controls (2N), for the peripheral
region 97 % did not differ (all proteins except for methylglutaconyl-CoA hydratase, row 18) whereas for the central region, 69 % did not (column 3, proteins in bold and italic). Also most of the proteins with a different expression between the 2N and 4N groups did not change expression with age in BD subjects. Together, these findings are a strong indication that especially the peripheral region of area 17 still resembles that of younger normal controls due to a BD-induced delay in maturation.
Table 1 List of 36 differentially expressed proteins in area 17

| Spot number | Process | Protein name                                                | NORMAL age | BD  | 2BD/2N | 4BD/4N |
|-------------|---------|-------------------------------------------------------------|------------|-----|--------|--------|
| 1           | o       | CRMP1                                                       | ↓↓         |     | -      | -      |
| 2           | o       | CRMP2                                                       | ↓↓         |     | -      | -      |
| 3           | o       | CRMP4                                                       | ↑↑         |     | -      | -      |
| 4           | o       | tubulin alpha-6 chain                                       | ↓↓         |     | -      | -      |
| 5           | o       | dynactin subunit 2                                          | ↑↓         |     | -      | -      |
| 6           | c       | Hsc70                                                       | ↑↑         |     | -      | -      |
| 7           | c       | endophilin-B2                                                | ↑↓         |     | -      | -      |
| 8           | m       | ATP synthase subunit beta                                   | ↑↓         |     | -      | -      |
| 9           | s       | septin 5                                                    | ↑↑         |     | -      | -      |
| 10          | m       | ATP synthase subunit d                                      | ↑↑         |     | -      | -      |
| 11          | m       | isocitrate dehydrogenase [NAD] subunit alpha, mitochondrial | ↑↑         |     | -      | -      |
| 12          | m       | heat shock 70 kDa protein 12A                                | ↑↓         |     | -      | -      |
| 13          | m       | MAGUK p55 subfamily member 2                                | ↓↓         |     | ↑↓     | ↑↓     |
| 14          | m       | alpha-1-syntrophin                                          | ↓↓         |     | ↑↓     | ↑↓     |
| 15          | m       | methylglutaconyl-CoA hydratase                               | ↑↑         |     | -      | -      |
| 16          | m       | peroxiredoxin-6                                             | ↑↑         |     | -      | -      |
| 17          | m       | glutathione S-transferase P                                 | ↑↑         |     | -      | -      |
| 18          | m       | Hsp90alpha isoform 2                                         | ↑↑         |     | ↑↑     | +      |
| 19          | m       | NFL                                                         | ↑↑         |     | -      | -      |
| 20          | m       | alpha-synuclein                                             | ↑↑         |     | -      | -      |
| 21          | m       | L-lactate dehydrogenase B chain                              | ↓↓         |     | ↑↓     | ↑↓     |
| 22          | m       | hnrNPL                                                      | ↓↓         |     | ↑↓     | +      |
| 23          | m       | glucose-6-phosphate isomerase                                | ↑↑         |     | -      | -      |
| 24          | m       | ATP synthase subunit alpha                                   | ↑↑         |     | -      | -      |
| 25          | m       | cyclophilin D                                               | ↑↑         |     | -      | -      |
| 26          | m       | glutamate dehydrogenase                                      | +          |     | +      | +      |
| 27          | m       | cytochrome b-c1 complex subunit 7                           | ↑↑         |     | +      | +      |
| 28          | m       | cytochrome c                                                | ↑↑         |     | +      | +      |
| 29          | m       | aldehyde dehydrogenase                                      | ↑↑         |     | -      | -      |
| 30          | m       | hnrNPH                                                      | ↑↑         |     | -      | -      |
| 31          | m       | hemoglobin subunit alpha                                     | ↑↑         |     | +      | +      |
| 32          | m       | hemoglobin subunit beta                                      | ↑↑         |     | +      | +      |
| 33          | m       | beta-synuclein                                              | ↑↑         |     | +      | +      |

Categorization was done according to the effect of age, BD, cortical region and the process that a particular protein is involved in. The first two 'age-regulated' columns visualize the presence and direction of difference (up or down) between the 2N and 4N group within central and peripheral area 17. The next two columns show the effect of age when reared under BD conditions, indicating the presence and direction of difference between the 2BD and 4BD group. In the next four columns, proteins were classified as '2BD-regulated' or '4BD-regulated' when their expression differed between a particular BD group and its age-matched normal control, or when there was a difference between the central and peripheral region in a BD group but not in its age-matched normal control group (+ or vice versa –). Between 4BD and 2N kittens, the peripheral region was molecularly similar for 97% (except protein in row 18), the central region was similar for 69% (25 proteins, marked in bold italics) and all other proteins showed higher expression in 4BD as compared with 2N. For CRMP2 and CRMP4 spot numbers that do not contain other proteins are in bold. Abbreviations: C: central; P: peripheral; o: outgrowth; s: synaptic transmission; c: clathrin-mediated endocytosis; m: metabolism; mR: mRNA metabolism and transport; ↑ – upregulation, or ↓ – downregulation. All presented differences are significant, p < 0.05.
Classification of identified proteins according to biological function

To reveal molecular pathways and biological functions potentially involved in cortical maturation, IPA software was applied to categorize the data set of the 36 identified proteins. The first two essential canonical pathways were ‘Parkinson’s Signalling’, containing synuclein alpha, septin 5, cytochrome c (p = 5.7E-04); and ‘Mitochondrial Dysfunction’, including the proteins involved in energy production, such as ATP synthase subunit alpha and beta, cytochrome c and cytochrome b-c1 complex subunit 7 (p = 7.07E-04). A significant canonical pathway was also ‘Semaphorin Signaling in neurons’ regulating neurite outgrowth, where CRMP1, CRMP2 and CRMP4 are the main players (p = 4.25E-03). Another canonical pathway was ‘Clathrin-mediated Endocytosis Signaling’ involving Hsc70, endophilin-B2 and albumin (p = 9.56E-02). Albumin was also a member of ‘Caveolar-mediated Endocytosis Signaling’ canonical pathway, which involved coatamer subunit epsilon isoform 4 (p = 1.15E-01). In the nervous system, albumin is a cargo protein transported to astrocytes not via clathrin- but caveolar-mediated endocytosis to promote synthesis and release of the neurotrophic factor oleic acid and subsequent neuronal differentiation and outgrowth [21]. Albumin was therefore not considered as a protein involved in clathrin-mediated endocytosis, but as a molecule transported via caveolar-mediated endocytosis.

Based on the IPA output we plotted a scheme presenting interactions between the identified proteins and their involvement in different biological processes. Figure 2a shows the interactions between 27 proteins related to three biological processes: outgrowth, clathrin-mediated endocytosis and synaptic transmission. Figure 2b illustrates a separate pathway linking the 11 proteins important for energy production. As eight out of the eleven identified mitochondrial energy metabolism-associated proteins had a higher expression in the 2BD group as compared to age-matched normal controls (Table 1), their upregulation suggests a higher energy demand to support neuronal activity and thus corresponds well with the previously revealed hyperactivity in area 17 of 2BD subjects [8].

The effect of age and BD on developmental profiles of proteins involved in outgrowth, synaptic transmission and clathrin-mediated endocytosis

To validate the 2-D DIGE results and IPA interpretation, we performed Western analysis for 4 proteins with a significant change in expression and belonging to the main biological processes highlighted in Fig. 2a: CRMP2 and CRMP4 for outgrowth, alpha-synuclein for synaptic transmission and Hsc70 for clathrin-mediated endocytosis. For synaptic transmission we additionally analyzed GAD65 and GAD67 expression in all experimental conditions as markers of inhibitory transmission.

Outgrowth

2-D DIGE revealed a diverse effect of age and BD on the different CRMP2 isoforms (Fig. 3a). Some isoforms increase, others decrease with age in normal but not in BD kittens. When probing for the overall CRMP2 expression level by Western analysis low levels were detected in 1N and 2N groups, which increased to the adult level by the age of 4 months (Fig. 3b). This age-dependent increase in CRMP2 expression was also observed in early onset BD animals but lasted longer, until 6 months of age (Fig. 3a, b). Nevertheless, in comparison to age-matched normal controls the total amount of CRMP2 did not differ for either early or late onset BD kittens (Fig. 3a, b).

In contrast to CRMP2, all CRMP4 isoforms showed a similar age dependent profile in normal and BD kittens as visualized by 2-D DIGE (Fig. 3c). Western blotting also revealed this initial decrease in expression until month 4, and an increase towards the age of 6 months, to achieve adult levels (Fig. 3d). In all BD conditions CRMP4 levels were higher as compared to age-matched normal controls in both regions of area 17 (Fig. 3c, d; Table 2). Nevertheless the expression profile remains parallel to the normal developmental course.

Synaptic transmission

GAD67 produces the main cellular pool of GABA, whereas GAD65 produces GABA that is preferentially packed into vesicles for fast neurotransmitter use [22–29]. The expression of GAD65 and GAD67 increased during normal cortical development achieving mature levels at 6 months of age for GAD67 and at 4 months for GAD65 for both central and peripheral area 17 (Fig. 4a, b). In early and late onset BD animals GAD67 expression levels did not differ from age-matched normal controls or across regions (Fig. 4a). The only difference we detected relates to the lack of a significant increase in GAD67 levels between month 4 and 6. The developmental increase of GAD65 expression was slower in BD compared to normal animals in both regions of area 17, achieving the highest level only in 6BD kittens (Fig. 4b). Additionally, in the central region of 4BD kittens GAD65 did not show a developmental increase, maintaining a lower level as compared to the 4N group (Fig. 4b). As such, the GAD65 expression level in 4BD was similar to that in 2N kittens (Fig. 4b). In 2N2BD, GAD65 expression did not differ from age-matched 4N controls or younger 2N animals. However, similar to 4BD, 2N2BD showed a lower GAD65 level than 6BD in both regions, indicating that both late onset and short early
Proposed schematic overview of protein interactions. Interactions among detected proteins and proteins described in literature that are involved in processes with regard to the development of primary area 17: outgrowth, synaptic transmission, clathrin-mediated endocytosis (a) and metabolism (b). Interactions are accompanied by numbers, referring to corresponding references: 1.[85] 2.[86] 3.[87] 4.[88] 5.[49] 6.[89] 7.[90] 8.[91] 9.[72] 10.[92] 11.[93] 12.[94] 13.[69] 14.[95] 15.[7] 16.[96] 17.[32] 18.[97] 19.[98] 20.[99] 21.[100] 22.[101] 23.[102] 24.[103] 25.[104] 26.[105] 27.[21] 28.[25] Grey rectangles – processes, rectangles with rounded corners – differential identified proteins; rectangles with rounded corners and dotted background – proteins studied by WB; proteins in an oval frame and dotted background – proteins studied by WB only; arrows – stimulation; line – interaction; perpendicular lines – inhibition.
Fig. 3 CRMP2 and CRMP4 protein expression analysis. Visualization and analysis of the relevant spots from the 2-D DIGE experiments containing CRMP2 (a) and CRMP4 (c); semi-quantitative Western blotting for CRMP2 (b) and CRMP4 (d). Overall, the CRMP2 expression increased with age and was not different in BD animals as compared to age-matched normal controls. The level of CRMP4 in normal controls decreased towards month 4 and again increased at the age of 6 months, and was higher in all BD animals as compared to age-matched normal controls. Asterisks above bars denote significant differences ($P < 0.05$) for a given region between age groups of a given condition (normal or BD). Numbers above BD-related bars denote the % statistical difference between BD and age-matched normal control groups ($P < 0.05$). Results are means with ± SD. Abbreviations: C – central, P – peripheral.
onset BD arrests the developmental increase in GAD65 levels (Fig. 4b).

Alpha-synuclein is involved in the regulation of vesicle storage and turnover as well as in the maturation and modulation of synaptic function [30–32]. It promotes SNARE-complex assembly during exocytosis in presynaptic terminals [32]. The Western blot experiment showed that specifically at the age of 1 month alpha-synuclein expression is lower in peripheral area 17. This centro-peripheral gradient is lost from 2N onwards. After the age of 6N, both cortical regions exhibit a drop to adult expression levels (Fig. 5). Western analysis confirmed the 2-D DIGE result showing a lower level of alpha-synuclein in the central region of 2BD kittens as compared to age-matched normal controls (Fig. 5). In 2N2BD subjects both central and peripheral area 17 showed a lower alpha-synuclein level as compared to age-matched normal controls, exhibiting a similar effect to that observed for early onset, age-matched 4BD kittens (Fig. 5).

**Clathrin-mediated endocytosis**

Western analysis for Hsc70 revealed an opposite expression profile for early onset BD kittens compared with normal animals (Fig. 6). In normal kittens, a decrease from the age of 2 months into adulthood characterized both regions of area 17, and in the 1 N group Hsc70 expression was lower in central as compared to peripheral area 17 (Fig. 6). In 2N2BD subjects both central and peripheral area 17 showed a lower alpha-synuclein level as compared to age-matched normal controls, exhibiting a similar effect to that observed for early onset, age-matched 4BD kittens (Fig. 5).

**Discussion**

Normal development of primary visual cortex is shaped by visual experience and mirrored by region-specific activity reporter gene expression, in conjunction with the previously described central-to-peripheral maturation gradients in the visual system (reviewed in [11]). During normal development, in cat, the peripheral region of area 17 still undergoes intensive developmental changes between the 2nd and 4th month of age when its central counterpart is already in a more mature state [8]. We also witnessed such a centro-peripheral maturation gradient in 1N and 2N kittens based on protein expression patterns. As predicted, BD exerted a cortical region-specific effect on these protein expression profiles. Several of these developmental protein expression changes occurring between 2 and 4 months in normal animals (Table 1; column normal/age; rows 1–20) were absent in BD subjects. In fact, protein expression in peripheral area 17 hardly differed between 4BD kittens and the younger 2N animals, confirming our previous results where we showed that BD exerts a stronger delay effect on the maturation of peripheral area 17 as measured by visually-induced activity reporter gene zif268 expression [8]. Additionally, under BD, centro-peripheral differences were observed up to 4 months. Together, these observations indicate that BD may not only delay but also prolong and enhance the centro-peripheral protein expression gradient related to the development of area 17. In sum, by exploiting the impact of BD on cortical maturation we could implicate four biological processes and thirty-six proteins in visual cortex development.

**Metabolism**

The constant energy demand for protein synthesis needed for neurite outgrowth and the formation of
new connections was reflected by changes in protein expression related to energy metabolism. It has been suggested before that processes that underlie synaptic plasticity may rely on changes in energy metabolism and expression of genes and proteins involved in metabolic processes, which is often defined as ‘metabolic plasticity’ [33–37] (reviewed in [38]). According to our 2-D DIGE data, 2 months of BD from birth, but not 4 months, induces a wide range of changes in expression of proteins involved in energy production, indicating the presence of a period of enhanced visual cortex plasticity. 2BD animals may thus exhibit more plasticity potential as compared to 2N animals, which are indeed already in the declining phase of the critical period for ocular dominance plasticity [39–41]. Indeed, in 1N kittens expression of mitochondrial genes was higher [42]. This hypothesis is in line with recent work [8], where we showed that 2BD kittens exhibited a pattern of high molecular activity in area 17, similar to 1-month-old controls, which are at the peak of the critical period for ocular dominance plasticity [40, 43]. Dark rearing also induces an upregulation of several mitochondrial genes (ATPase 6, cytochrome b, NADH dehydrogenase subunit 4 and 2; [42]). Altogether, these observations indicate that the high expression of proteins involved in energy metabolism may not only reflect the neuronal activity level and metabolic demand, but also is an indicator of neuronal plasticity. The upregulation of L-lactate dehydrogenase in 2BD kittens supports this interpretation, as L-lactate signaling was shown to be involved in plasticity processes during memory formation where it mediates molecular changes such as induction of phospho-CREB, Arc and phospho-cofilin [44]. Furthermore, L-lactate derived from astrocytes also affects NMDA receptor signaling and induces expression of IEGs including arc, zif268 and c-fos in vitro and in vivo, inducers of synaptic plasticity and activity [45].

Outgrowth

The differential regulation of CRMP2 and CRMP4 under BD may be related to their cell-type specific expression and to the signaling pathways related to structural plasticity they are involved in. CRMP2 mediates repulsive semaphorin3A (Sema3A) signaling through the Rac-dependent pathway [46, 47] or a Rho kinase-dependent cascade reaction. Growth cone collapse is induced by activation of RhoA GTPase and Rho kinase (ROCK), leading to subsequent microtubule disassembly (reviewed in [48]). CRMP4 also interacts with RhoA GTPase [49], but is not known to be a ROCK substrate. While both CRMP2 and CRMP4 are capable of binding to tubulin heterodimers [50], CRMP4 is suggested to interact mainly with the actin...
cytoskeleton as it was shown to bundle with F-actin [51]. These data suggest that BD may reorganize the actin cytoskeleton in cat area 17 in a way that slows down the developmentally regulated formation of new connections.

We observed an age-dependent decrease in CRMP4 expression in both regions of area 17 as previously observed for cats and rodents [1, 2, 52, 53]. Crucial to the interpretation of our data, CRMP2 and CRMP4 exhibit cell-type specific expression. CRMP4-immunoreactive neurons appear mainly as non-pyramidal and parvalbumin-positive, a marker for a distinct subset of inhibitory interneurons, whereas CRMP2-immunopositive neurons are mainly parvalbumin-negative and display a clear pyramidal shape, typical for excitatory neurons [3, 54]. The development of inhibitory connectivity in cat area 17, measured by the number of symmetric synapses, follows a linear increase that reaches adult values by month 4 [55, 56], when the lowest CRMP4 expression is observed. A correlation between the CRMP4 expression profile and the development of inhibitory connectivity would be in line with the specific immuno-localization of CRMP4 in parvalbumin-positive interneurons [3].

**Synaptic transmission**

Consistent with previous results by Guo and coworkers [5], in animals with normal visual experience, GAD65 and GAD67 achieved adult expression levels at 4 and 6 months of age respectively, in line with observations in humans [57]. In BD animals we observed a delay in the
developmental increase of GAD65. In mice with a reduced synthesis of GABA, due to knockout of the gene encoding GAD65, ocular dominance plasticity is not present and the critical period is postponed until an appropriate level of inhibition is experimentally acquired [58, 59]. Experience-dependent plasticity therefore requires a minimal level of cortical inhibition to establish a threshold in the excitation/inhibition balance (for review see [60, 61]). The delay in the developmental increase in GAD65 in BD kittens thus reflects the delayed maturation of area 17. Dark rearing was shown to decrease the number of GAD65 puncta on layer 2/3 pyramidal neurons in mouse visual cortex [62, 63]. In rat, immature GABAergic inhibition in the visual cortex has also been observed in the context of dark-rearing from birth [64]. Together these observations indicate that visual experience is permissive to reach mature inhibitory networks. In the 2N2BD group no difference in GAD expression was detected when compared to age-matched normal controls. It is not surprising since a late-onset dark rearing period, preceded by normal visual experience during the first 3 weeks of life, also did not change GABAergic transmission in 5-week-old dark reared rats [64].

The expression of alpha-synuclein, a modulator of synaptic neurotransmission, was higher in kittens than in adult cats. Interestingly, for alpha-synuclein we observed higher levels in central as compared to peripheral area 17 in 1-month-old normal control kittens. A Zebra Finch homolog of alpha-synuclein is expressed in the song control nucleus when song plasticity is at its maximum [65]. Thus, the higher expression of alpha-
synuclein in the central region at 1N may be an indicator of an earlier critical period onset for central area 17 as compared to peripheral area 17 [66]. Alpha-synuclein was shown to promote exocytosis. Its decreased expression in both regions of 2BD and in the central region of 4BD kittens, in combination with our observations for GAD65 and CRMP4, may suggest a negative regulation of inhibitory neurotransmission [30, 32].

Clathrin-mediated endocytosis
In our study a higher level of Hsc70 is observed in kittens than in adult cats, which is in line with a higher intensity of receptor recycling in younger animals [67]. High levels of Hsc70 may be related to an increased intensity of clathrin-mediated endocytosis responsible for developmental rearrangements in N-methyl-D-aspartate receptor (NMDAR) subunit (NR1, NR2A, NR2B) number and composition at the excitatory glutamate synapse (Fig. 2a, [68]). In kittens, all NMDAR subunits show a peak expression at 2 months of age with a subsequent decline into adulthood [7]. Much of the decline in expression of NMDAR subunits occurs between the age of 2 and 4 months in cat [6, 7]. Enhanced internalization of NMDARs at early developmental stages contributes to the preferential insertion of NR2B over NR2A, while the decline in NMDAR internalization during neuronal maturation may be related to the stabilization of the mature NR2A/NR2B expression ratio at the synapse [68, 69] (reviewed in [70]).

BD resulted in a downregulation of Hsc70 in both area 17 regions in 2BD and in the peripheral region only of 4BD as compared to age-matched normal controls, in agreement with our previous observation that retinal lesion-induced visual cortex plasticity involves downregulation of Hsc70 [37]. Our 2-D DIGE data showed that the other protein indicated by IPA to be involved in clathrin-mediated endocytosis, endophilin-B2 [71], was also downregulated under BD, in the central region of 2BD kittens. Another protein that was shown to enhance clathrin-mediated endocytosis after neuronal stimulation is alpha-synuclein [72], which reveals a comparable modulation by BD to that observed for Hsc70.

Hsc70 may be involved in the formation of a complex of proteins that anchor GAD65 to the synaptic vesicle membrane [25, 29]. Thus, the decrease of Hsc70 expression up to the age of 4 months under early onset BD may not only indicate a downregulation of clathrin-mediated endocytosis but also a decrease in the activity of synaptic vesicles, reduced GABA synthesis and neurotransmission. Clathrin-mediated endocytosis is a major pathway for synaptic vesicle recycling (reviewed in [73]), thus downregulation of this pathway is again in line with downregulation of inhibitory synaptic transmission.

Distinct behavioral outcomes of pattern deprivation in adult cats can be foreseen in the developmental molecular profiles
Overall, our assessment of the developmental cortical proteome confirms the recently described immature state of the primary visual cortex due to early onset BD [8]. For 2BD kittens we detected an upregulation of proteins involved in energy production as if the cortex is in a more immature state. 4BD kittens exhibited a delay in maturation specifically for peripheral area 17. Exactly the peripheral region of 4BD subjects had a proteomic signature of more immature 2-month-old normal control kittens and correspondingly, centro-peripheral protein expression gradients were detected up to 4 months of BD. This molecular immaturity of the primary visual cortex at the end of a given BD period seems to permit a different behavioral outcome in relation to the functional compensation induced by ensuing normal visual input [16]. Once adult, 2BD cats indeed outscore normal cats on most of the tested motion perception tasks. 4BD cats perform at the same level as normal subjects. In contrast, delayed onset 2N2BD or 6 BD cats show specific deficiencies in motion perception tasks [16], here mirrored by an adult-like molecular profile at the end of the BD period (summarized in Table 2), as described for zif268 expression [8]. Specifically these two BD regimes did not induce a downregulation of Hsc70 (Table 2), a protein involved in clathrin-mediated endocytosis. 6BD subjects also did not show a downregulation of alpha-synuclein, a protein involved in synaptic transmission (Table 2) and show the most profound anatomical deficiencies in adulthood specific to motion perception, e.g., anatomical rearrangements of motion sensitive alpha retinal ganglion cells [14] and severe impairment of the simplest motion detection task [17 compare with 16, 18]. Most likely the impaired development of primary visual cortex becomes stabilized at some point, thereby preventing further modifications even in the context of restoration of normal visual experience.

Conclusions
Taking advantage of the recently described impact of a postnatal BD period on primary visual cortex maturation [8] we could implicate four biological processes and thirty-six proteins in subregion-specific cortical development within area 17. We could correlate transient negative regulation of neurite outgrowth, synaptic transmission and clathrin-mediated endocytosis to the previously reported differences in behavioral outcome in adult cats with regard to the timing and duration of such a BD period in early life [16]. Combined, our findings suggest that the cortical plasticity potential to functionally recover from an early BD period once normal visual input is restored may evolve as a function of
the specific sets of protein expression changes or delays, instigated by the loss of pattern vision in early life.

Methods

Animals

All experiments were carried out in accordance with the European Parliament and the Council Directive of September 22nd 2010 (2010/63/EU). The cats were raised under a daily photoperiod of 12 h light and 12 h darkness with water and food ad libitum (Nencki Institute, Warsaw, Poland). All efforts were made to minimize animal discomfort. The cat brain material for this protein investigation has already been used in parallel in another study [8].

To screen for proteome changes in relation to area 17 maturation under normal visual stimulation (N) and to screen for the effect of binocular deprivation from patterned visual experience (BD), we analyzed central and peripheral regions of area 17 of cats with normal visual experience at the age of two (2N, n = 3) and four (4N, n = 3) months and of cats binocularly deprived for either two (2BD, n = 3) or four (4BD, n = 3) months from eye opening (P8). Western blot analysis was also performed on additional animal groups: normal kittens of one (1N, n = 2) and six months (6N, n = 3), normal adult cats of 1–2 years (n = 3), kittens binocularly deprived for six months from eye opening (6BD, n = 3) and kittens from a delayed onset BD group that were deprived for the third and fourth month of life after two initial months of normal visual input (2N2BD, n = 3).

BD was always achieved by having the cats wearing double thickness linen masks covering their eyes. This procedure reduces retinal illumination to a similar level as lid suturing, but is less traumatic [74]. The masks were replaced daily in a normally lit animal facility room where the kittens lived. The changing procedure lasted no longer than one minute per day for each cat, which is not sufficient to maintain normal vision [75, 76] and allowed constant adjustment of the size of the masks to the growing head. The masks were removed at the end of the deprivation period.

Prior to administration of an overdose of sodium pentobarbital (Nembutal, 60 mg/kg, i.p.) all animals were maintained overnight in total darkness followed by 1-h light stimulation. Brains were dissected, instantly frozen by immersion in dry ice cooled isopentane (Poch, Gliwice, Poland) and stored at −80 °C.

Protein extraction

We collected tissue from the central and peripheral region of area 17 at Horsley-Clarke level posterior 6.0–7.0. Using the in situ hybridization films for zf268 [8] and the cat visual cortex map of [77] as a guide, we punched ~10 mm² cortical tissue, containing all cortical layers (Fig. 1a), from 3–4 consecutive 200 μm-thick frontal sections for each experimental and normal control condition to obtain sufficient protein material.

For 2-D DIGE, brain tissue was transferred to 100 μl ice-cold lysis buffer, containing 7 M urea (Chem-Lab), 2 M thiourea (Fluka), 4 % w/v CHAPS (Applichem), 1 % w/v dithiothreitol (DTT) (Applichem), 40 mM Tris base (Sigma), and Complete Protease Inhibitor Cocktail (Roche Diagnostics). Brain tissue was homogenized on ice, briefly centrifuged at 13 000 rpm, sonicated, followed by a complete solubilization of the proteins for 1 h at RT. The proteins were sonicated again and centrifuged for 20 min at 13 000 rpm at 4 °C to precipitate cell debris. The supernatant was dialyzed against milli-Q water for 2 h to remove residual salt using a membrane with a 500 Da cut-off (Spectra/Per, Biotech, Omnilabo). Protein concentrations were determined according to the Qubit™ Quantitation Platform (Invitrogen) using a Qubit® fluorometer (Invitrogen, Merelbeke, Belgium). Samples were kept at −80 °C.

For Western blotting, the collected brain tissue was homogenized in 100 μl lysis solution (2 % w/v sodium dodecyl sulfate (SDS) [Sigma- Aldrich], 50 mM Tris–HCl [Sigma-Aldrich], 10% glycerol [Acros Organics], pH 6.8) containing 4 μl protease inhibitor (Complete Protease Inhibitor Cocktail tablets; Roche Diagnostics). After mechanical homogenization by drill-driven pestles, all the samples were sonicated for 5 x 10 s, heated at 70 °C for 5 min, and centrifuged for 20 min at 13 000 rpm at 4 °C. The supernatant was isolated and the total protein concentration was determined as described above.

Two-dimensional difference Gel electrophoresis

The fluorescent cyanine dyes, Cy2 (Cy2), propyl-Cy3 (Cy3), and methyl-Cy5 (Cy5) were in-house synthesized [1, 2] according to the method by Ünlü et al. [78]. All other chemicals were purchased from GE Healthcare, unless mentioned otherwise. Pre-cast Immobiline DryStrips (24 cm, pH 3–11 nonlinear) were rehydrated overnight in DeStreak Rehydration Solution containing 0.5 % v/v immobilized pH gradient (IPG) buffer in a reswelling tray covered with paraffin oil (Merck). The next day 50 μg protein of each cat area 17 sample was randomly labeled with either propyl-Cy3 or methyl-Cy5. Equal fractions of all samples were pooled and 50 μg of this pool was labeled with Cy2 to serve as an internal standard. The minimal amount of dye that gave a maximum number of spots and the highest signal-to-noise ratio was set to approximately 200 pmol as described previously [1, 2]. The samples were incubated for 30 min on ice in the dark during the labeling process; subsequently, the reaction was terminated by addition of 1 μl lysine (10 mM; Merck) for 15 min. The Cy2-, Cy3-, and Cy5-labeled fractions were mixed together, and an equal
volume of lysis solution was added. Isoelectric focusing (IEF) was performed on an Ettan IPGphor Cup Loading Manifold system according to manufacturer's instructions. Actual run conditions were 300 V for 3 h, 600 V for 3 h, followed by a 6-h gradient to 1000 V, a 3-h gradient to 8000 V, and 8 h at 8000 V for a total of 50 kVh (at 50 μA/strip). After IEF, the strips were equilibrated twice for 15 min in equilibration buffer (6 M urea, 34.5 % v/v glycerol and 10 % w/v SDS in Tris–HCl buffer [1.5 M, pH 8.8]). DTT (1 % w/v) was added to the first equilibration step and 4.5 % w/v iodoacetamide (Sigma-Aldrich) to the second step. Electrophoresis of the IPG strips was done on 1.5-mm-thick SDS-polyacrylamide gels (12.5 % T; 2.6 % C) in the Ettan DALT twelve system for 30 min at 30 mA and 24 h at 13 °C at 15 mA/gel.

2-D DIGE gel image analysis and statistics
Gels were scanned with the Ettan DIGE Imager (software 1.0; GE Healthcare) and generated gel image triplets (Cy2, Cy3, and Cy5) comprising the CyDye-labeled proteins. Quantitative analysis was carried out with the DeCyder 2D difference analysis software (Version 7.0; GE Healthcare). Spot detection and matching was performed automatically with the DeCyder Batch processor. The gel-to-gel matching was also checked manually followed by statistical analysis of protein abundance change between samples in the biological variation analysis (BVA) module embedded in the DeCyder Software [37, 79, 80]. Spots of interest differentially expressed at least in two comparisons with p < 0.05, and spots differentially expressed in one comparison with p < 0.01 were further analyzed with MS (39 spots fulfilled the criteria, Fig. 1b).

Protein identification
Two preparative gels were run under the same conditions as described above with the exception of the first dimension separation, which was run for a total of 50 kVh (at 50 μA/strip). Each gel was loaded with 1 mg of protein from the pool sample, from which only a 50-μg fraction was labeled with Cy3. Glass plates were pretreated with BindSilane, and 2 reference markers were applied to enable automatic spot picking. The preparative gels were scanned in the Ettan DIGE Imager to obtain an image of the Cy3 signal.

Subsequently, the total protein load was visualized by Lava Purple total protein fluorescent stain according to manufacturer’s instructions (Fluorotechnics), and the gels were scanned again. Matching with the analytical gels was as before carried out automatically with manual correction by the BVA module of the DeCyder software. A pick list of the proteins of interest was generated and imported into the Spot Picker Version 1.20 software that controls the Ettan Spot Picker (GE Healthcare).

Protein identification
In collaboration with the Centre de Recherche Public-Gabriel Lippmann in Luxemburg, MS and MS/MS spectra were acquired using a 5800 MALDI TOF-TOF (AbSciex, Sunnyvale, CA, U.S.A.) and calibrated using the 4700 peptide mass calibration kit (Applied Biosystems). Proteins were identified by searching against the NCBI database, limited to the taxonomy Mammalia (downloaded on 4/06/2012, 1,063,527 sequences), using an in-house MASCOT server (version 2.3.0 Matrix Science, www.matrixscience.com, London, U.K.). All searches were carried out defining trypsin as cleavage agent and allowing for 2 missed cleavages. A mass window of 100 ppm was tolerated for the precursor mass and 0.75 Da for fragment ion masses. The search parameters allowed for carboxymethylation of cysteine as fixed modification and oxidation of methionine and tryptophan (double oxidation, and kynurenin formation) as variable modifications. Proteins were considered as identified when two, none overlapping, individual peptides surpassed the peptide score threshold or when the expect value <5.0e-005. When this criterion was not met, additional precursors were selected and searched using the above-described parameters. In total, we identified 36 unique proteins (Additional file 1: Table S1) in 32 spots (all 39 spots classified for identification with MS are indicated in Fig. 1b). Occasionally, proteins with the same name were detected in more than one spot due to the fact that the 2-D DIGE method is sensitive enough to separate different isoforms and posttranslational modifications of one protein (Table 1, e.g., rows 2 and 3; CRMP2 and CRMP4). In one case, Western blot results were opposite to the 2-D DIGE findings, likely due to the co-occurrence of two proteins, Hsc70 and CRMP2, in one spot (nr 623). Therefore only the Western blot data were used to describe the Hsc70 results.

The 36 identified proteins from the 2-D DIGE analysis were analyzed by means of the QIAGEN’s Ingenuity Pathway Analysis (IPA®, QIAGEN Redwood City, www.qiagen.com/ingenuity) software. IPA calculates the significance value of a given canonical pathway as the probability that the pathway is associated with the data set by random chance. To reveal key biological pathways, functions and molecular networks comprising the identified proteins we applied the stringent Benjamin-Hochberg (B-H) multiple testing correction method.

Western blotting and statistics
To obtain the optimal protein load, a protein dilution series was performed with total protein amounts of 0.5-25 μg per lane. A concentration that resulted in a good signal to noise ratio and still was in the linear range of the detection system was chosen. For CRMP2, CRMP4, Hsc70, alpha-synuclein, GAD65/67 this resulted in 2.5,
8, 4, 30 and 15 μg respectively. After the addition of 5 μl reducing agent (10X, Invitrogen) and 2 μl LDS sample buffer (4X, Invitrogen), the samples were denatured (10 min, 70 °C). Protein samples were loaded onto a 4-12 % Bis-Tris NuPage gel (Invitrogen). Electrophoresis was carried out using the Xcell4 SureLock Midi-Cell module (Invitrogen) according to the manufacturer’s instructions and subsequently transferred to a PVDF or Nitrocellulose (for alpha-synuclein) membrane (iBlot, Gel Transfer Stack; Invitrogen). The Spectra Multicolor High range protein ladder (ThermoScientific) was used as molecular weight standard. After blotting, the membrane was incubated for 1–2 h in a 5% ECL blocking agent (GE Healthcare, Buckinghamshire, UK) in Tris-saline (0.01 M Tris, 0.9% NaCl, 0.1% TX-100, pH 7.6) and incubated overnight with a primary antibody (diluted in 5% blocking agent in Tris Saline) against CRM2 (1/24000, mouse Ab, generously provided by Dr. Y. Ihara, University of Tokyo, Department of Neuropathology, Japan), CRM4 (1/10000, rabbit Ab, AB5454 Millipore), Hsc70 (1/2000, rabbit Ab, SPA-816, Stressgen Bioreagents), alpha-synuclein (1/1000, rabbit Ab, in house produced and characterized by Prof. Veerle Baekelandt, KU Leuven, Department of Neurosciences, Belgium [81]), GAD65/67 (1/6000, rabbit Ab, ab11070, Abcam). The next day, the blots were washed in Tris-Saline (3x5min) and 30 min incubated with a horseradish peroxidase-conjugated secondary goat anti-mouse (GaM-HRP, 1/50000, Dako, Glostrup, Denmark), goat anti-rabbit (GaR-HRP, 1/50000; Dako, Glostrup, Denmark) or donkey anti-goat Ab (1/50000, sc-2020, Santa Cruz Biotechnology) (diluted in 5% blocking agent in Tris Saline), followed by a rinse in Tris-Saline (3x5min) and Tris-stock (1x5min; 0.05 M Tris, pH 7.6). Immunoreactivity was visualized using chemiluminescence detection (Supersignal West Dura, Thermo Scientific, Pierce) on ECL hyperfilm (GE Healthcare). The protein bands were semi-quantitatively evaluated by densitometry (ImageQuant TL v. 7.0; GE Healthcare). For the Hsc70 antibody two bands were detected in samples of young cats (up to 4 months), independent of visual manipulation, as previously reported for adult cats with retinal lesions [37]. Only the lower one (MW of 73 kDa, [82]) was considered to be specific according to the molecular-weight size marker.

To account for intra-gel and inter-gel variability including loading differences, incomplete transfer, or position on the blot, a total protein stain (LavaPurple, Gelcompany) was used rather than the use of a single reference protein [37, 83] (Additional file 2: Figure S1, Additional file 3: Figure S2, Additional file 4: Figure S3, Additional file 5: Figure S4) according to manufacturer’s instructions. For semi-quantitative densitometry, non-uniform staining was corrected for by inter-lane measurements and normalization. For each protein of interest, the specific protein band per cat was normalized to its corresponding normalized total protein stain. Also, each experiment on the same cat was repeated two times. A reference sample (pool) consisting of a mixture of each prepared tissue sample was run with the same optimal amount of protein on each gel to gauge blot-to-blot variability (first left lane in each blot in Figs. 3, 4, 5 and 6). Statistical analysis of WB data was performed using a nested-design ANOVA model to investigate the effects of group, cortical region and blot nested in cat [84] by means of data analysis software system STATISTICA version 10. There was a significant effect of interaction between group and region for GAD65 (p = 0.0269, F = 0.24, df = 8), alpha-synuclein (p = 0.0001, F = 14.01, df = 8), Hsc70 (p = 0.0001, F = 6.305, df = 8). There was a significant effect of group for CRM2 (p = 0.0056, F = 3.57, df = 6), CRM4 (p = 0.0001, F = 68.60, df = 6) and GAD67 (p = 0.0001, F = 8.45, df = 6). A post hoc test was carried out using the Tukey HSD method. Statistical differences were indicated for p < 0.05.

Additional files

Additional file 1: Table S1. List of proteins identified by mass spectrometry. (PDF 104 kb)
Additional file 2: Figure S1. Total protein stain for normalization of CRMPs specific bands. (PDF 607 kb)
Additional file 3: Figure S2. Total protein stain for normalization of GADs specific bands. (PDF 331 kb)
Additional file 4: Figure S3. Total protein stain for normalization of α-synuclein specific bands. (PDF 316 kb)
Additional file 5: Figure S4. Total protein stain for normalization of Hsc70 specific bands. (PDF 262 kb)

Abbreviations

2-D: Two-Dimensional; BD: binocular deprivation; C: central; c: clathrin-mediated endocytosis; CRM: collapsin response mediator protein; DIGE: Difference Gel Electrophoresis; GAD: Glutamic acid decarboxylase; Hsc70: Heat shock cognate 70; IEF: Isoelectric focusing; IPA: Ingenuity Pathway Analysis; LGN: lateral geniculate nucleus; m: energy metabolism; mR: mRNA metabolism and transport; MS: mass spectometry; N: normal; NMDA: N-methyl-D-aspartate receptor; α: outgrowth; P: peripheral; ROCK: RhoA GTPase and Rho kinase; s: synaptic transmission; WB: Western blot.

Competing interests

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be constructed as a potential conflict of interest.

Authors’ contributions

All authors had full access to all the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis. Study concept and design: KB, LA. Acquisition of data: KL-M, T-TH. Analysis and interpretation of data: KL-M, JK, JN. Drafting of the manuscript: KL-M, JN. Critical revision of the manuscript for important intellectual content: JN, MK, KB, LA. Statistical analysis: KL-M and KB. Administrative, technical and material support: MZ, AVP. Study supervision: KB, LA. All authors read and approved the final manuscript.
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Author details
1Laboratory of Neuroplasticity, Nencki Institute of Experimental Biology, 02-093 Warsaw, Poland. 2Laboratory of Neuroplasticity and Neuroproteomics, KU Leuven - University of Leuven, 3000 Leuven, Belgium. 3Laboratory for Neurobiology and Gene Therapy, KU Leuven - University of Leuven, 3000 Leuven, Belgium.

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