Acta Physiologica Hungarica, Volume 102 (2), pp. 216–227 (2015)

DOI: 10.1556/036.102.2015.2.12

Altered mitochondrial response to activation of T-cells in neonate

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> Received: September 19, 2014 Accepted after revision: March 12, 2015

Mitochondrial functions have a major impact on T-cell functionality. In this study we characterized whether mitochondrial function in the neonatal T-cells differs from that in the adult T-cells during short T-cell activation. *Methods:* We used flow cytometry methods to test mitochondrial mass and to monitor mitochondrial Ca²⁺ levels, mitochondrial potential and superoxide generation in parallel with cytoplasmic Ca²⁺ levels during phythohaemagglutinine-induced activation of CD4+ and CD8+ T-cells of 12 term neonates and 11 healthy adults. *Results:* Baseline mitochondrial mass of CD4+ and CD8+ cells was lower in the neonate than in the adult. In comparison with the adult, neonatal resting CD4+ T-cells had lower cytoplasmic Ca²⁺ levels and this was associated with normal activation induced Ca²⁺-response. During short-term activation cytoplasmic Ca²⁺-response was lower in neonatal than in adult CD8+ T-cells. Mitochondrial Ca²⁺ uptake was increased in CD4+ neonatal T cells while it decreased in CD8+ T-cells. Mitochondrial depolarization was increased in CD4+ and decreased in CD8+ neonatal T-cells compared to adults. Superoxide generation was higher and equal in neonatal CD4+ and CD8+ cells, respectively, compared to the adult ones. *Conclusion:* Our data suggest that neonatal T-cells exhibit marked differences in mitochondrial function and superoxide generation compared to adult T-cells.

Keywords: calcium, flow cytometry, mitochondrial membrane potential, neonate, superoxide, T-cell

Clinical experience and experimental data support that neonatal T-cells are immunologically less competent than those in the adult (1). Upon activation the production of inflammatory cytokines in CD3+ T-cells is lower in the neonate than in the adult (7, 13). The major factors explaining the reduced response of neonatal T-cells is probably the absence of pre-existing immunological memory (2) and increased number of CD71+ erythroid cells (5). However, we have also observed that the intracellular Ca²⁺-response of the neonatal T-cells to activation also presents specific characteristics (23).

Proper T-cell activation is a result of coordinated work of intracellular signaling machinery (6). As a result of T-cell receptor (TCR) stimulation, cytoplasmic Ca^{2+} level ($[Ca^{2+}]_c$) is transiently increased and it activates Ca^{2+} -dependent signaling pathways (8). We have found that Ca^{2+} -response during short-term activation of T-cells is lower in the neonate than in the adult. The immaturity of potassium channels on neonatal cell membrane may be an explanation for this finding (23).

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Now we focused on mitochondrion, another central player of T-cell activation. Changes in $[Ca^{2+}]_c$ are associated with alterations in mitochondrial Ca^{2+} levels ($[Ca^{2+}]_m$) (4). Mitochondria have an important role in T-cell activation as they modulate the Ca^{2+} -signaling during TCR induced activation. Additionally mitochondrial Ca^{2+} -signal regulates energy metabolism and has also a major impact on other specific T-cell functions (19).

Appropriate generation of reactive oxygen species (ROS) is a fundamental element of T-cell activation (10, 22). ROS modulates signaling pathways and transcription of several genes (25). TCR-signaling induces mitochondrial O_2^- generation in a Ca^{2+} dependent manner (3). Phagocyte-type NADPH-oxidase and mitochondria are the major sources of ROS generation during the first 15 minutes of T-cell activation (10).

In this study we compared mitochondrial mass and function including mitochondrial Ca²⁺ kinetics and mitochondrial membrane potential and ROS generation in CD4+ and CD8+ T cells of the neonate with those of the adult.

Materials and Methods

Patients

We used cord blood samples of twelve healthy term neonates [10 girls and 2 boys, gestational age: 39 (38–40) weeks, median (range), birth weight: 3375 (2510–4040) grams, median (range)] and peripheral blood samples of 11 healthy adults [3 women and 8 men, age: 25 (22–44) years, median (range)] (Table I).

	Number	Gender (F/M)	Birth weight (g)	Gestational age (neonate) or age	Ethnicity
Neonates	12	10/2	3375 (2510–4040)	39 (38–40) weeks	Caucasian (Hungarian)
Adults	11	3/8	N.A.	25 (22–44) years	Caucasian (Hungarian)

Table I. Clinical data and patient characteristics. Data are expressed as median (range)

Ethical statement

The study adhered to the tenets of Declaration of Helsinki. Written informed consent was obtained from the adults and the mothers of the neonates. An independent institutional ethical committee, Semmelweis University Regional and Institutional Committee of Science and Research Ethics, approved this study.

Solutions and reagents

Ficoll Paque; Amersham Biosciences AB, Uppsala, Sweden; phosphate buffered saline (PBS) (2 mM KH₂PO₄, 9.5 mM Na₂HPO₄.2H₂O, 136.7 mM NaCl); anti-human mAbs: anti-CD4 PE-Cy7 (cat. no. 557852), anti-CD8 APC-Cy7 (cat. no. 557834) (both from PharMingen, San Diego, CA, USA); Fluo-3-AM, Rhod2/AM, tetramethylrhodamine methylester (TMRM), MitoTracker Green and Dihydroethidium (DHE) dyes as well as Pluronic F-127 were purchased from Molecular Probes (Karlsbad, CA) and they were freshly dissolved in dimethyl sulfoxide (DMSO) or ethanol according to the manufacturer's protocol before the measurements. Phytohemagglutinin (PHA), ionomycin, carbonylcyanide-4-(trifluoromethoxy)-phenylhydrazone (FCCP), RPMI-1640 medium were purchased from Sigma–Aldrich (St. Louis, MO, USA).

Cell isolation

Peripheral blood mononuclear cells (PBMC) and cord blood mononuclear cells (CBMC) were separated by a standard density gradient centrifugation with Ficoll Paque (27 min, 400 g, 22 °C) from freshly drawn blood collected in lithium heparin-treated tubes (BD Vacutainer and BD Biosciences, San Jose, CA, USA). This cell suspension was washed twice in PBS. From then on, cells were kept throughout staining with fluorescent markers and measurement on flow cytometer in a modified RPMI medium. The Ca²⁺ concentration of the modified RPMI medium was set to 2 mM by addition of crystalline CaCl₂.

Surface staining

For surface marker staining, PBMCs and CBMCs were incubated with anti-CD4 PE-Cy7 and anti-CD8 APC-Cy7 anti-human conjugated mAbs according to the manufacturers' instructions.

Cell activation

At the beginning of each measurement 2 min baseline was recorded. T-cell activation was initiated by phytohemagglutinin (15 mg/ml in final concentration). In each experiment fluorescence emission of sequentially measured cells was monitored for 12 minutes, the time which is needed for the decline of the PHA induced calcium signal in adult and cord blood human T-cells (23). In average, about 1 x 10⁶ cells were measured.

Cytoplasmic Ca²⁺kinetics measurement

Cytoplasmic free Ca²⁺ level was detected by loading the cells with Fluo-3-AM (emission maximum, 526 nm, recorded with a 530/30 nm band pass filter (BP filter)). Cells were loaded with 2.6 mM Fluo-3-AM and 0.02% Pluronic F-127 for 30 minutes at 30 °C. Cells were washed once before measurements. At the end of the measurements ionomycin was added to the samples to verify the specificity of the signal (17).

Measurement of superoxide generation

Dihydroethidium (DHE), a redox-sensitive probe reacts with superoxide and results in the formation of a fluorescent dye (emission maximum, 605 nm, recorded with 610/20 nm BP filter). Cells were loaded with 1 mM DHE for 18 minutes at 37 °C (17).

Measurement of baseline mitochondrial mass and investigation of the kinetics of mitochondrial Ca²+ level, mitochondrial membrane potential upon activation

Mitochondrial mass was estimated by staining with potential insensitive mitochondrial dye, MitoTracker Green (emission maximum: 516 nm recorded in with a 530/30 nm BP filter). Cells were loaded with 150 nM MitoTracker Green for 30 minutes at 37 °C in the dark. Cells were washed once before measurements.

Mitochondrial Ca²⁺ level was monitored using Rhod2/AM (emission maximum, 581 nm, recorded with a 575/26 nm BP filter). Cells were loaded with 2.5 mM Rhod2/AM and 0.02% Pluronic F-127 for 20 minutes at 30 °C and washed once before measurements. At the end of each measurement FCCP was added to the sample to verify the specificity of the signal (17).

Mitochondrial membrane potential was estimated by staining with TMRM (emission maximum, 567 nm recorded with 575/26 nm BP filter) in quench/dequench mode. Cells were loaded with 1 μ M TMRM for 20 min at 37 °C in the dark and washed once before measurement. At the end of the measurements FCCP was added to the sample to verify the specificity of the signal.

Equipment and gating methods

All flow cytometric measurements were performed on a BD FACSAria flow cytometer (BD Biosciences, San Jose, CA) equipped with 488- and 633-nm excitation lasers and data were processed using the FACSDiVa software.

The population of lymphocytes was gated according to forward scatter characteristics and side scatter characteristics. PE-Cy7 conjugated anti-CD4 and APC-Cy7 conjugated anti-CD8 antibodies were used for the gating of CD4+ and CD8+ T-cell subpopulations. The gating strategy of flow cytometry data is shown in Fig. 1.

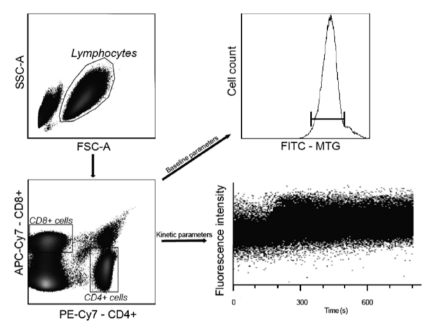


Fig 1. Staining and gating strategy. The population of lymphocytes was gated from PBMCs according to Forward Scatter Characteristics and Side Scatter Characteristics. The CD4+ or CD8+ subpopulations were differentiated according to the surface staining and used either for the measurement of baseline parameters (for example mitochondrial mass by MitoTracker Green (MTG)) or for kinetic measurements

Data analysis

Data acquired from the measurements were evaluated with specific software developed at our laboratory. The core of this software is an improved version of an algorithm (15) based on fitting of logistic or double logistic functions to each recording (14). The logistic function was used to characterize continuously increasing fluorescence values, while the double logistic function was used to describe measurements that have an increasing phase, a peak and a decreasing phase as time passes. The software also calculates parameter values describing each function, such as the maximum value (Max), the time to reach maximum value (t_{max}), slope at 50% of maximum (Slope), ending value (End) and the area under the curve (AUC). One unit (U) of the AUC value is defined as one relative intensity value in 1 s, where relative intensity values are the rate of actual intensity values divided by intensity values at 0 s.

Comparison of parameter distributions of the two groups was made by probability binning method (21), Mann–Whitney tests were performed as tests of normality (performed

according to Kolmogorov–Smirnoff) indicated non-normal distribution of data. Mann–Whitney tests were used to test the difference between baseline parameters in the different cell populations. Two tailed p-values obtained with less than 0.05, were considered as significant.

Results

Mitochondrial mass

To examine the mitochondrial mass in resting CD4+ and CD8+ T-cell populations isolated from newborn cord blood and adult peripheral-blood, we measured the whole cell fluorescent intensity of MitoTracker Green, a mitochondrial specific fluorescent dye. Mitochondrial mass was significantly decreased in the neonatal CD4+ T-cell populations compared to the adult CD4+ T-cells, while the difference in the CD8+ T-lymphocytes was not significant (Fig. 2A).

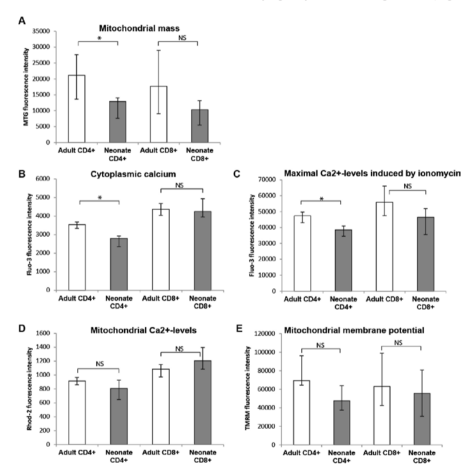


Fig. 2. Comparison of resting T-cell parameters. Median fluorescence values of the mitochondrial specific fluorescence dye MitoTracker Green (MTG) (A); baseline and ionomycin induced cytoplasmic Ca²⁺-levels measured by Fluo-3 (B and C respectively); mitochondrial Ca²⁺-levels estimated by Rhod-2 (D); mitochondrial membrane potential measured by TMRM (E) in adult and neonate CD4+ or CD8+ T-cell subpopulations. Abbreviations: A-CD4+ – adult CD4+ T-cells; A-CD8+ – adult CD8+ T-cells; N-CD4+ – neonatal CD4+ T-cells, N-CD8+ – neonatal CD8+ T-cells. Data are expressed as median and interquartile range.

Baseline cytoplasmic Ca^{2+} levels, mitochondrial Ca^{2+} levels, mitochondrial membrane potential

Lower cytoplasmic Ca²⁺ levels in neonatal CD4+ T-cells

In order to compare the cytoplasmic Ca²⁺ levels of unstimulated adult peripheral and neonatal cord blood CD4+ and CD8+ T-cells, T-cells were loaded with Fluo-3-AM, a calcium sensitive fluorescent dye. Baseline cytoplasmic Ca²⁺ levels of neonatal CD4+ T –cells were lower compared to adult CD4+ lymphocytes whereas no difference was observed between the 2 groups in CD8+ T-cells (Fig. 2B). The addition of the calcium ionophore ionomycin indicating that the capacity for pharmacologically induced calcium flux in neonatal CD4+ T cell was reduced compared to adult CD4+ T-cells, while it was comparable in neonatal and adult CD8+ T-cells (Fig. 2C).

Comparable basal mitochondrial Ca^{2+} levels and membrane potential in neonatal and adult CD4+ and CD8+ T-cells

When investigating the role of mitochondria in Ca²⁺-metabolism of neonatal and adult T-cells, first we measured basal mitochondrial Ca²⁺ levels. Mitochondrial Ca²⁺ levels of unstimulated neonatal and adult CD4+ or CD8+ T-cells were comparable (Fig. 2D). Mitochondrial membrane potentials of resting CD4+ or CD8+ T-cells were equivalent in the two analyzed groups (Fig. 2E).

PHA induced changes in cytoplasmic Ca^{2+} levels, mitochondrial Ca^{2+} levels, mitochondrial membrane potential and superoxide generation

Decreased cytoplasmic Ca²⁺ levels in neonatal CD8+ cells following PHA induced activation

We stimulated T-cells with PHA and monitored the initial fluctuation of cytoplasmic Ca²⁺ levels. AUC, Max or End and Slope parameter values were calculated from the fitted functions and compared between the two groups. During short-term activation cytoplasmic Ca²⁺ response (in term of parameters AUC and Slope) was lower in neonatal than in adult CD8+ T-cells (Fig. 3A and C) while no difference in neonatal and adult CD4+ T-cells was observed (Fig. 3A and B).

Elevated mitochondrial Ca²⁺ levels in neonatal CD4+following PHA induced activation In order to examine the effect of cytoplasmic Ca²⁺ signaling on mitochondrial Ca²⁺-uptake and mitochondrial function we measured mitochondrial Ca²⁺ levels and mitochondrial membrane potential after stimulation with PHA. The extent and kinetics of mitochondrial Ca²⁺-uptake induced by PHA addition (characterized by AUC, Slope and End) was increased in CD4+ T cells obtained from neonates compared to adults (Fig. 4A and B), while neonatal CD8+ cells contained less Ca²⁺ than adult CD8+ T-cells at the end of the measurement (End value) (Fig. 4A and C).

Increased mitochondrial depolarization in neonatal CD4+ and decreased in CD8+ T-cell following PHA stimulation

In parallel to mitochondrial calcium uptake, mitochondrial membrane potential showed similar alterations following PHA stimulation: increased depolarization (AUC and Max) in neonatal CD4+ T-cells compared to adult (Fig. 5A and C) and reduced depolarization in CD8+ neonatal T-lymphocytes compared to that one of adult (AUC, Slope and Max) (Fig. 5A and C).

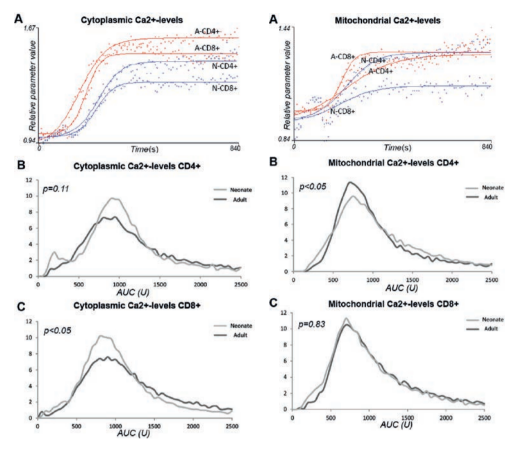


Fig. 3. PHA induced kinetic changes in cytoplasmic Ca²⁺ levels. Figure shows the calculated median functions (A) (records of representative samples). Histograms showing the distributions of Area Under the Curve (AUC) parameter of fitted functions in CD4+ (B) and CD8+(C) T-cells. Abbreviations: A-CD4+ – adult CD4+ T-cells; A-CD8+ – adult CD8+ T-cells; N-CD4+ – neonatal CD4+ T-cells, N-CD8+ – neonatal CD8+ T-cells

Fig. 4. PHA induced kinetic changes in mitochondrial Ca²⁺ levels. Figure shows the calculated median functions (A) (records of representative samples). Histograms showing the distributions of Area Under the Curve (AUC) parameter of fitted functions in CD4+ (B) and CD8+(C) T-cells. Abbreviations: A-CD4+ – adult CD4+ T-cells; A-CD8+ – adult CD8+ T-cells; N-CD4+ – neonatal CD4+ T-cells, N-CD8+ – neonatal CD8+ T-cells

Superoxide generation

To see whether these changes in mitochondrial calcium metabolism and function affect the mitochondrial superoxide generation, we measured the fluorescent intensity changes of DHE after PHA stimulation. Surprisingly superoxide generation was also higher (based on AUC parameter) in neonatal than in adult CD4+ T-cells (Fig. 6A and B) whereas no differences were observed between neonatal and adult CD8+ T-cells (Fig. 6A and C). Table II summarizes our results acquired by PHA stimulation of CBMC and PBMC T-cells.

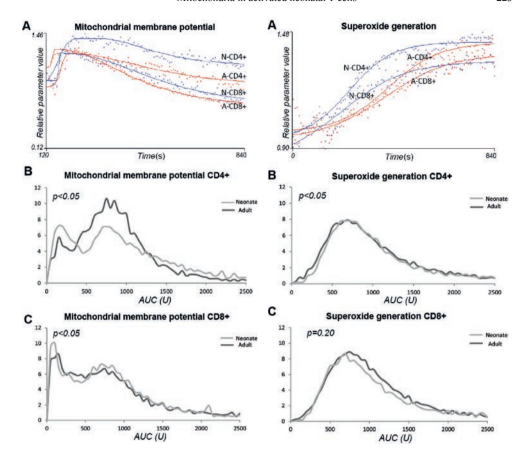


Fig. 5. PHA induced kinetic changes in mitochondrial membrane potential. Figure shows the calculated median functions (A) (records of representative samples).
 Histograms showing the distributions of Area Under the Curve (AUC) parameter of fitted functions in CD4+ (B) and CD8+(C) T-cells. Abbreviations: A-CD4+ – adult CD4+ T-cells; A-CD8+ – adult CD8+ T-cells; N-CD4+ – neonatal CD4+ T-cells, N-CD8+ – neonatal CD8+ T-cells

Fig. 6. PHA induced kinetic changes in superoxide generation. Figure shows the calculated median functions (A) (records of representative samples). Histograms showing the distributions of Area Under the Curve (AUC) parameter of fitted functions in CD4+ (B) and CD8+(C) T-cells. Abbreviations: A-CD4+ adult CD4+ T-cells; A-CD8+ adult CD8+ T-cells; N-CD4+ neonatal CD4+ T-cells, N-CD8+ neonatal CD8+ T-cells

Discussion

In this study, we have observed that in addition to earlier observations done on cytoplasmic Ca²⁺-response, neonatal T-cells present other functional characteristics distinct from T-cells of the adult (11, 23). Now we measured reduced baseline cytoplasmic Ca²⁺ levels and reduced Ca²⁺-influx capacity in CD4+ neonatal T-cells.

After these observations in Ca²⁺-handling we focused on mitochondrial function during T-cell activation. The rationale is the previous data suggesting that mitochondria are central players of T-cell activation and Ca²⁺-signaling (9).

Table II. Effects of PHA stimulation on cord blood and adult CD4+ and CD8+ T cell populations

Parameter	T-cell subpopulation							
	Adult CD4+ (n = 10)	Neonate CD4+ $(n = 11)$	Adult CD8+ $(n = 10)$	Neonate CD8+ $(n = 11)$				
Cytoplasmic Ca ²⁺ -levels (CD4+-double logistic+ function; CD8+ logistic+ function)								
AUC	1193[799–2263]	1059[789–1760]	1155[791–1881]	982 ^b [726–1445]				
Slope	0.0046[0.0016-0.0177]	0.0030[0.0014-0.0136]	0.0022[0.0010-0.0080]	0.0015 ^b [0.0006–0.0073]				
Max or End	1.62[1.06-3.26]	1.39[1.02-2.53]	1.62[1.16–2.58]	1.32[0.96–2.01]				
Mitochondrial Ca ²⁺ -levels (logistic+ function)								
AUC	939[691–1541]	989a[693-1582]	960[687–1497]	899[645–1377]				
Slope	0.0009[0.0004-0.0041]	0.0018a[0.0009-0.0043]	0.0008[0.0004-0.0060]	0.0024 ^b [0.0007-0.0260]				
End	1.37[1.02–2.19]	1.46 ^a [0.94-2.43]	1.46[1.04–2.11]	1.19 ^b [0.85–1.79]				
Mitochondrial membrane potential (double logistic+ function)								
AUC	797[518–1101]	880°[465–1507]	773[358–1415]	767 ^b [357–1270]				
Slope	0.0245[0.0029-0.2203]	0.0061[0.0010-0.0255]	0.0326[0.0037-0.1707]	0.0289 ^b [0.0036-0.1460]				
Max	1.15[0.78–1.53]	1.27°[0.69-2.13]	1.13[0.52-2.01]	1.10 ^b [0.60–1.77]				
Superoxide generation (O ₂ ⁻) (logistic+ function)								
AUC	1001[634–1909]	1058a[676-2412]	955[652–1474]	1008[656–1960]				
Slope	0.0010[0.0006-0.0023]	0.0022[0.0007-0.0082]	0.0008[0.0005-0.0013]	0.0029[0.0008-0.0175]				
End	1.51[0.99–2.89]	1.49[0.96–3.68]	1.36[0.94–2.06]	1.39[0.89–2.94]				

The measured parameter values of logistic+ and double logistic+ functions were AUC (area under the curve in units [U]), Max (double logistic+ function – maximum value in relative parameter value [rpv]), Slope (slope at the first 50% value of Max), and End (logistic+ function – ending value). Data are expressed as median and interquartile range.

Indeed we measured lower mitochondrial mass in neonatal CD4-cells than in those from the adult. Previous study has shown that CD8+ memory T cells contain a higher amount of mitochondria compared to naïve T-cells (24). Neonates, compared to adults, have higher ratio of CD45RA+ naïve T-cells (~90% and ~55%, respectively). The mitochondrial mass of CD4+ naïve and memory T cells has not been investigated so far. Although we did not distinguish naïve and memory T-cells in our experiments, the higher ratio of naïve T-cells might contribute to the lower mitochondrial mass of neonatal CD4+ and CD8+ T-cells.

The mass of mitochondria has a major effect on intracellular Ca²⁺-handling. Mitochondria act not only as Ca²⁺-buffers but they have a major impact on Ca²⁺-channel regulation (20). Mitochondrial Ca²⁺-uptake reduces cytoplasmic Ca²⁺peaks, but delays the return of [Ca²⁺]_c to the baseline. This feature was strikingly different in CD4+ and CD8+ T-cell populations. In the presence of lower mass of mitochondria, neonatal CD4+ T-cells took up more, while CD8+ cells took up less Ca²⁺ than those in the adult. These altered mitochondrial Ca²⁺-handling characteristics might shape the cytosolic Ca²⁺-signal leading to an earlier decline of cytosolic Ca²⁺-signal. As there are no data available concerning the mitochondrial Ca²⁺-handling mechanism in neonatal T-cells further studies are required to explain these

^a Neonatal cord blood samples were compared with adult samples (CD4+ T-cell population), p < 0.05

^b Neonatal cord blood samples were compared with adult samples (CD8+ T-cell population), p < 0.05

differences. This issue however is of major importance for immune modulation, as the diminished cytoplasmic Ca²⁺levels were reported to be responsible for the reduced NFAT-associated gene expression (9, 13).

Of note, mitochondrial depolarization in T-cells upon activation is associated with the altered mitochondrial Ca²⁺-uptake in the neonatal T cells. Increased mitochondrial depolarization upon PHA induced T-cell activation was detected in neonatal CD4+ cells, as a consequence of elevated mitochondrial Ca²⁺-sequestration. On the other hand, mitochondrial depolarization was lower in neonatal CD8+ T-cells characterized by smaller Ca²⁺-uptake. These findings suggest that the difference in mitochondrial depolarization of T-cells between the neonate and the adult is attributable, at least partly to different Ca²⁺-handling.

Mitochondrial calcium uptake has a major impact on ROS generation. Activation of T-cells is accompanied by increased reactive oxygen species generation (in a Ca²⁺-dependent manner (3)). ROS facilitate the expression of the oxidation-dependent nuclear transcription factors NF-κB and AP-1 in T-cells (18). In spite of the diminished CD4+ T-cell mitochondrial mass in the neonatal T-cell, we observed increased ROS production in these cells. This finding suggests altered regulation of mitochondrial ROS production in the very early phase of cord blood T-cell activation and/or the dysfunctionality of intracellular antioxidant defense mechanisms. Additionally, a previous work has demonstrated that between 15 and 75 minutes of T-cell activation in the neonate, these cells produce increased amount of ROS after chemical stimulation (ionomycin + phorbol-dibutyrate) (16) and decreased amount after anti-CD28 mAb stimulation. We propose that the elevated initial and then decreased ROS generation contribute to the different activation characteristics of neonatal T-cells. However, we have to note some limitations regarding our superoxide measurements done in a period when flow cytometry technique was a widely accepted approach for the detection of superoxide generation. However, today HPLC based methods are recommended to detect 2-hydroxyethidium (2-OH-E+), the fluorescence product formed from the DHE/O2•reaction (12). Further studies applying these techniques are required to confirm our results about superoxide generation in T-cells; however, our observations provide a sound basis for these experiments.

Upon evaluation of our findings, we are to consider the limitations of our experiments. Number of samples measured and blood amount (25 ml) of adult and newborn groups were limited, and the study was performed *in vitro*.

Conclusions

Our data suggest that neonatal T-cells present different characteristics during short-term activation. In addition to reduced cytoplasmic Ca^{2+} -signaling the mitochondrial mass is decreased in neonatal CD4+ T-cells. We measured elevated mitochondrial Ca^{2+} -sequestration and mitochondrial depolarization in CBMC T-cells. All these alterations might contribute to the elevated O_2^- production and ROS signaling (Fig. 7). These findings indicate that intracellular machinery in its complexity presents different features between neonatal and adult T-cells that may contribute to the immaturity of adaptive immunity of the neonate.

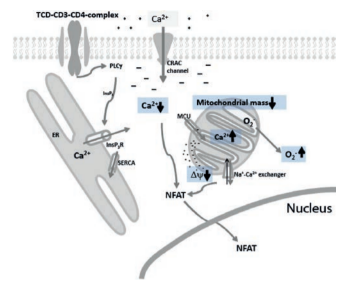


Fig. 7. Schematic representation of neonatal CD4+ T-cell activation.

Schematic representation of CD4+ T-cell activation showing the parameters altered in neonate compared to adult. In addition to reduced cytoplasmic Ca²⁺-signaling the mitochondrial mass is decreased in neonatal CD4+ T-cells. Decreased mitochondrial mass is responsible for the elevated mitochondrial Ca²⁺-sequestration and mitochondrial depolarization which contribute to the elevated O₂ production. All these changes might contribute to reduced NFAT-associated gene expression in neonatal CD4+ T-cells

Acknowledgement

This work was supported by OTKA Grants 84087 and 101661.

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