Monitoring the expression of purinoceptors and nucleotide-metabolizing ecto-enzymes with antibodies directed against proteins in native conformation

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Abstract Following their release from cells, ATP and NAD, the universal currencies of energy metabolism, function as extracellular signalling molecules. Mammalian cells express numerous purinoceptors, i.e., the nucleotide-gated P2X ion channels and the G-protein-coupled P2Y receptors. Signalling through purinoceptors is controlled by nucleotide-metabolizing ecto-enzymes, which regulate the availability of extracellular nucleotides. These enzymes include ecto-nucleoside triphosphate diphosphohydrolases (ENTPD, CD39 family) and ecto-nucleotide pyrophosphatase/phosphodiesterases (ENPP, CD203 family). Investigation of these receptors and enzymes has been hampered by the lack of available antibodies, especially ones that recognize these proteins in their native conformation. This study reports the use of genetic immunization to generate such antibodies against P2X1, P2X4, P2X7, ENTPD1, ENPTD2, ENPTD5, ENPTD6, ENPP2, ENPP3, ENPP4, ENPP5, and ENPP6. Genetic immunization ensures expression of the native protein by the cells of the immunized animal and yields antibodies directed against proteins in native conformation (ADAPINCs). Such antibodies are especially useful for immunofluorescence and immunoprecipitation analyses, whereas antibodies against synthetic peptides usually function well only in Western-blot analyses. Here we illustrate the utility of the new antibodies to monitor the cell surface expression of and to purify some key players of purinergic signalling.

Keywords Monoclonal antibodies · Purinoceptor · Ecto-nucleotidases · Genetic immunization

Abbreviations

ADAPINCs Antibodies directed against proteins in native conformation
AEBSF 4-(2-Aminoethyl) benzenesulfonyl fluoride hydrochloride
ATP Adenosine triphosphate
ENPP Ecto-nucleotide pyrophosphatase/phosphodiesterase
ENTPD Ecto-nucleoside triphosphate diphosphohydrolase
NAD Nicotinamide adenine dinucleotide
PO Peroxidase
PVDF Polyvinylidene difluoride

Introduction

Following their release from cells, the universal currencies of energy metabolism, ATP and NAD, function as extracellular signalling molecules [1–5]. Molecular cloning has identified 7 nucleotide-gated ion channels (P2X purinoceptors) and >8 nucleotide-activated G-protein-coupled receptors (P2Y purinoceptors) [6–10]. Signaling through
purinoceptors is effectively controlled by nucleotide-metabolizing ecto-enzymes, which regulate the availability of extracellular nucleotides [11]. Prominent roles are played by members of two families of ecto-enzymes: the ecto-nucleoside triphosphate diphosphohydrolases (ENTPD, CD39 family) [12] and the ecto-nucleotide pyrophosphatase/phosphodiesterases (ENPP, CD203 family) [13].

The mammalian P2X family encompasses seven members. All P2X purinoceptors contain cytosolic N- and C-terminal tails, two membrane spanning domains, and an extracellular domain of approximately 280 amino acids containing the ligand-binding site and 10 conserved cysteine residues that likely form five intrachain disulfide bonds [14, 15]. The extracellular domains of the respective mouse and human P2X orthologues show high (ca. 80–90%) sequence identity; the paralogues within a given species show lower (40–50%) identity.

The mammalian ENTPD family encompasses eight members [16]. ENTPD1–4, 7 and 8 have a structure similar to that of P2X purinoceptors, i.e., cytosolic N- and C-terminal tails, two transmembrane spanning domains, and an extracellular domain of approximately 440 amino acids, encompassing the ligand-binding site and four conserved cysteines likely engaged in intrachain disulfide bridges [12]. ENTPD1–3 and 8 are expressed as cell surface enzymes, ENTPD4 and ENTPD7 as luminal enzymes of intracellular organelles. ENTPDases share structural similarities with the actin/HSP70/sugar kinase superfamily [12]. ENTPD5 and ENTPD6 lack the second transmembrane domain. Moreover, in ENTPD5, the region corresponding to the first transmembrane domain of the other family members is N-terminal and functions as a signal peptide, resulting in the secretion of ENTPD5 into the lumen of the endoplasmic reticulum. ENTPD6 is a type II transmembrane protein. ENTPD orthologues show high (ca. 80–90%) sequence identity; the paralogues within a given species show lower (30–50%) identity.

The ENPP family encompasses seven members [13]. ENPP1–3 are type II transmembrane proteins with a structure analogous to that of ENTPD6, i.e., an N-terminal cytosolic tail, a single transmembrane domain and an extracellular catalytic domain of approximately 400 amino acids. ENPP4–7, in contrast, are type I membrane proteins. A hydrophobic N-terminus acts as a signal peptide, the extracellular catalytic domain is followed by a single transmembrane domain and a short C-terminal cytosolic tail. ENPPs 4–7 consist of catalytic domains only, the catalytic domains of ENPPs 1–3 are flanked by genetically fused upstream somatomedin B and downstream nuclease-like domains [13]. ENPPs share structural similarities with the alkaline-phosphatase superfamily [17, 18]. The catalytic domain of ENPP orthologues shows high (ca. 80–90%) sequence identity; the paralogues within a given species show lower (40–50%) identity.

Despite their relevance for purinergic signalling, investigation of these receptors and enzymes has been hampered by the lack of available antibodies, especially ones that recognize the enzymes in their native conformation. Antibodies raised by immunization with synthetic peptides derived from the known amino acid sequence of a protein of interest generally work well in Western-blot analyses but often fail to recognize the native protein on the cell surface (Fig. 1). Such antibodies can be used successfully for monitoring the overall expression level of the protein of interest in a population of cells but cannot be used to assess cell surface expression on individual cells. Genetic immunization, in contrast, ensures that the protein of interest is expressed in native conformation by the cells of the

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**Fig. 1** DNA immunization vs. peptide immunization.
immunized animal and yields antibodies directed against proteins in native conformation (ADAPINCs) [19, 20]. Such antibodies function well in immunofluorescence, FACS, ELISA, and immunoprecipitation analyses, i.e., assays in which antipeptide antibodies often fail. We have previously employed this technique to generate highly specific polyclonal and monoclonal antibodies against ecto-ADP-ribosyltransferases and against murine P2X7 [20–23]. Here we report the generation and characterization of similar antibodies against other key players of purinergic signalling and illustrate the utility of these antibodies for monitoring the cell surface expression of their cognate antigens.

**Materials and methods**

Cloning of expression vectors and cell transfections

Clones encoding full-length open-reading frames representing members of the P2X, ENTPD, and ENPP families were purchased from the mammalian genome collection (www.rzpd.de). Clones that were not already in an expression vector (pCMV-SPORT6) were subcloned into the pCDNA6 expression vector. Further expression constructs were kindly provided by James Wiley, Sydney (human P2Xγ), Richard J. Evans, Leicester (human P2X1), François Rassendren, Montpellier (mouse P2X4), Ruth Murrell-Lagnado, Cambridge (rat P2X4), and Astrid Kehlen, Halle (hENPP2). An expression construct for nuclear green fluorescent protein was generated by cloning the DNA-binding domain of the LKLF transcription factor [24] into a C-terminal fusion protein into the pEGFP-N1 vector (Clontech). Expression constructs were transfected into CHO and HEK cells (5 × 10^6 cells in 200 μl PBS i.v.) at room temperature. Beads were washed twice in PBS/1% Triton X-100 by incubation on a roller for 30 min at room temperature. Beads were washed twice in PBS/1%

CHO cells were co-transfected with expression constructs for purinoceptors or ecto-enzymes and nuclear GFP (5 and 0.5 μg/10^6 cells, respectively). Cells were transferred onto 96-well plates (4 × 10^5 cells/well) 6 h post-transfection. Then, 20 h after transfection, cells were fixed for 10 min in 2% paraformaldehyde (PFA) and then stained with serial dilutions of immune serum for 30 min at room temperature. To detect ENTPD5, which is not bound to the cell surface, cells were fixed in 4% PFA and subsequently permeabilized with 0.5% IGEPAL (Sigma) before staining. Bound antibody was detected with PE-conjugated antirat or antirabbit IgG (1:100) (Dianova). Cell nuclei were counterstained with Hoechst 33342. Cells were analyzed with a Zeiss Axiovert 25 microscope equipped with the Canon PowerShot G2 digital camera and appropriate filters for visualizing Hoechst stain (Zeiss filter set 01), green fluorescent protein (Zeiss filter set 10) and PE fluorochrome (Zeiss filter set 15). Photographs were taken at 40× magnification with fixed camera settings (digital zoom: 4.3×, aperture: 5.0, exposure time: 2 s for GFP and PE, 1/8 s for Hoechst 33342). GFP and PE images were merged using Adobe Photoshop software.

FACS analyses

For FACS analyses, cells were harvested by brief trypsinization (5 min at room temperature) and washed in serum-containing medium. Cells (5 × 10^5/100 μl) were stained either directly with Alexa-488-conjugated antibodies (1 μg) or with serially diluted immune serum followed by PE-conjugated antirat or antirabbit IgG (1:100, Dianova). Gating was performed on living cells on the basis of propidium iodide exclusion.

Immunoprecipitation analyses

Antibodies were conjugated to Protein G Sepharose beads (Pharmacia) (1 μg antibody/20 μL beads in 200 μL PBS containing 1% Triton X-100) by incubation on a roller for 30 min at room temperature. Beads were washed twice in PBS/1%
Triton X-100 before use for immunoprecipitation. Cells (10^7/ml) were lysed in PBS, 1% Triton-X100, 1 mM AEBSF (Sigma) for 20 min at 4°C. Insoluble material was pelleted by centrifugation, and the cleared lysates (2×10^7 cells in 200 μL) were subjected to immunoprecipitation using 20 μL antibody-conjugated beads by incubation on a roller for 30 min at room temperature. Beads were washed four times in PBS/1% Triton X-100. Proteins were eluted from the beads by incubation in SDS-PAGE sample buffer for 15 min at 70°C. Eluted proteins were size-fractionated on precast SDS-PAGE gels (Invitrogen) and blotted onto PVDF membranes. Purinoceptors and ecto-enzymes were detected with serially diluted serum followed by peroxidase (PO)-conjugated antirabbit IgG (1:5,000) using the ECL system (Amersham).

Immunoblot analyses

Cells (10^7/ml) were pelleted by centrifugation and lysed by resuspension in SDS-PAGE sample buffer (Invitrogen). DNA was sheared by brief ultrasonication on ice (two 15-s pulses with an MSE ultrasonicator at maximum power setting). Insoluble material was pelleted by centrifugation, and soluble proteins (5×10^5 cell equivalents/lane) were size-fractionated on precast SDS-PAGE gels (Invitrogen) under reducing conditions and blotted onto PVDF membranes. Purinoceptors and ecto-enzymes were detected with serially diluted serum followed by PO-conjugated antirabbit IgG (1:5,000) using the ECL system (Amersham).

Results and discussion

Use of transiently transfected CHO cells to assess the specificity and titres of antisera against purinoceptors and ecto-enzymes

Antibodies raised by genetic immunization of rabbits and rats with cDNA expression constructs for purinoceptors, ENTPDs, and ENPPs were assayed for reactivity and specificity by indirect immunofluorescence staining of CHO cells, 24 h after cotransfection with a construct encoding nuclear GFP and the cDNA expression construct used for immunization (Fig. 2 and Table 1). By this time, transfected cells were clearly distinguishable from untransfected cells by virtue of their green fluorescent nuclei (Fig. 2). Specific antibodies were detected on the basis of bright cell surface staining of transfected cells and lack of reactivity with untransfected cells. Antipeptide antisera against the same antigens, in contrast, did not show any detectable reactivity in this assay (Table 1).

Monitoring cell surface expression of purinoceptors and ecto-enzymes by flow cytometry

The utility of the ADAPINC antisera for monitoring cell surface expression was assessed further by flow cytometry (FACS analyses) of transiently transfected HEK cells (Fig. 3). The results confirm that the antisera react...
specifically with appropriately transfected cells and that they do not react with untransfected HEK cells. Importantly, the antisera recognize the respective purinoceptor/ecto-enzyme on the cell surface of living cells. In contrast, antipeptide antisera against the same antigens did not show any detectable reactivity with the transfected cells in these assays. Further, the use of the ADAPINC antisera for FACS analyses permits an assessment of the relative expression level of the cognate protein on the cell surface of individual cells as well as an assessment of the number of transfected vs. untransfected cells in the cell population. These analyses...
parameters cannot be quantified appropriately by Western-blot analyses, since for these analyses, lysates are prepared from cell populations. We further used the FACS-based assay system to test whether the antisera cross-react with other members (paralogues) of the receptor/enzyme family and/or the same receptor/enzyme (orthologue) in other species (human, rat, or mouse). The results reveal that most antisera are specific for the receptor/enzyme used for immunization and do not cross-react with other members of the same protein family. Some antisera, however, did react with the orthologous protein, in particular at low dilutions, e.g., antiserum CR30 raised against mouse P2X\textsubscript{1} shows some reactivity with the human P2X\textsubscript{1} orthologue (Fig. 3a).

Some of the antisera recognize the denatured cognate antigen in Western-blot analyses

We next tested the utility of the ADAPINC antisera for monitoring overall protein expression levels in immunoblot analyses (Fig. 4). To this end, we size-fractionated the proteins from lysates of untransfected and of transiently transfected HEK cells by SDS-PAGE followed by immunoblot analyses. The results show that only some of the ADAPINC antisera detected bands of the expected size. For comparison, we again performed parallel analyses with antisera raised against the same proteins by peptide immunization. The results show that the antipeptide sera generally detected bands of the expected size. The C-terminal peptide is commonly used for peptide immunization. In the case of P2X\textsubscript{7}, this sequence is highly conserved between rat, mouse and human orthologues, and the corresponding antipeptide serum recognized the respective orthologues (not shown). In contrast, in the case of P2X\textsubscript{4}, the mouse and human

![Figure 4](image1)

**Fig. 4** Immunoblot analyses of purinoceptor (a) and ecto-enzyme (b) expression by transfected HEK cells. Untransfected (u) or HEK cells transfected with mouse (m) or human (h) purinoceptors or ecto-enzymes were solubilized 24 h post-transfection with SDS-PAGE sample buffer. Proteins in cell lysates were size-fractionated by SDS-PAGE and subjected to immunoblot analyses using the indicated antisera. Bound antibodies were detected with peroxidase-conjugated secondary antibody and the ECL system

![Figure 5](image2)

**Fig. 5** Immunoprecipitation of purinoceptors by antibodies raised via genetic immunization. Untransfected (u) and transfected (t) HEK cells were solubilized 24 h post-transfection with PBS/1% Triton X-100. Purinoceptors were immunoprecipitated from cell lysates by incubation with the indicated antisera bound to Protein G Sepharose. Precipitates were subjected to immunoblot analysis as in Fig. 4, using the indicated antipeptide antisera for detection
sequences differ in two and five positions, respectively, from the rat peptide used for immunization, and this antipeptide serum shows only weak cross-reactivity with these orthologues (Fig. 4a, Ab α-mP2X4).

Affinity purification of purinoceptors and ecto-enzymes

Finally, we tested the utility of the ADAPINC antisera for immunoprecipitation of the cognate proteins from cell lysates (Fig. 5). To this end, we prepared lysates of untransfected and of transiently transfected HEK using the nondenaturing nonionic detergent Triton-X-100. Lysates were incubated with ADAPINC antibodies immobilized on protein-G sepharose beads, and proteins bound to washed beads were subsequently analyzed by SDS-PAGE followed by immunoblot analyses with antipeptide antibodies. The results show that all of the ADAPINC antisera efficiently precipitated proteins of the expected size from cell lysates. This underscores the utility of these antibodies for purifying native receptors and ecto-enzymes from lysed cells.

Conclusion

Using genetic immunization, we have raised highly specific polyclonal and monoclonal antibodies against key players of purinergic signalling, i.e., P2X1, P2X4, and P2X7 purinoceptors and ENTPD1, ENPTD2, ENPTD5, ENPTD6, ENPP2, ENPP3, ENPP4, ENPP5, and ENPP6 enzymes. Our findings underscore the utility of genetic immunization for generating highly specific polyclonal and monoclonal antibodies directed against proteins in native conformation (ADAPINCs) [19, 20]. These antibodies are valuable tools for assessing the expression levels of the native protein by immunofluorescence analyses and flow cytometry, i.e., assays in which antipeptide antibodies often fail. The antibodies described here provide useful tools for further characterization of the structure and function of these purinoceptors and ecto-enzymes.

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Conflict of interest statement The Haag and Koch-Nolte labs receive a share of the revenues from commercial sales of the antibodies described here by Analytical Services North, Hamburg.

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