Abstract. Mouse and human macrophages express a plasma membrane receptor for extracellular ATP named P2Z/P2X\textsubscript{7}. This molecule, recently cloned, is endowed with the intriguing property of forming an aqueous pore that allows transmembrane fluxes of hydrophobic molecules of molecular weight below 900. The physiological function of this receptor is unknown. In a previous study we reported experiments suggesting that the P2Z/P2X\textsubscript{7} receptor is involved in the formation of macrophage-derived multinucleated giant cells (MGCs; Falzoni, S., M. Munerati, D. Ferrari, S. Spisani, S. Moretti, and F. Di Virgilio. 1995. J. Clin. Invest. 95:1207–1216). We have selected several clones of mouse J774 macrophages that are characterized by either high or low expression of the P2Z/P2X\textsubscript{7} receptor and named these clones P2Z\textsubscript{hyper} or P2Z\textsubscript{hypo}, respectively. P2Z\textsubscript{hyper}, but not P2Z\textsubscript{hypo}, cells grown to confluence in culture spontaneously fuse to form MGCs. As previously shown for human macrophages, fusion is inhibited by the P2Z/P2X\textsubscript{7} blocker oxidized ATP. MGCs die shortly after fusion through a dramatic process of cytoplasmic septionation followed by fragmentation. These observations support our previous hypothesis that the P2Z/P2X\textsubscript{7} receptor is involved in macrophage fusion.

Purinergic P2X receptors are emerging as one of the most interesting new families of plasma membrane receptors recently described. Molecular cloning has shown that they are formed by subunits possessing only two probable transmembrane domains, with both the carboxy and amino termini on the cytoplasmic side of the plasma membrane and a central (280 amino acids) extracellular domain, rich in cysteine residues (Brake et al., 1994; Valera et al., 1994; Surprenant et al., 1996). Other plasma membrane receptor families sharing in part this structural motif are the inward rectifying K\textsuperscript{+} channel (Kir) of insulin-secreting cells, the amiloride-sensitive Na\textsuperscript{+} channel of epithelial cells, the mechanosensitive channel of Escherichia coli, and the mechanosensitive channels of Caenorhabditis elegans (deg-1, mec-4, and mec-10; Corey and Garcia-Anoveros, 1996; North, 1996). Among these, P2X and Kir are the only members that are known to be controlled by a soluble ligand, ATP in the case of P2X and ADP in that of Kir (Surprenant et al., 1995; Nichols et al., 1996).

P2Z/P2X\textsubscript{7}, the largest receptor/channel of the P2X subfamily (595 amino acids), differs from other members of the P2X subfamily by the presence of a long cytoplasmic carboxy tail that is essential for the pore-forming activity, as elegantly shown by Surprenant et al. (1996). Although P2Z/P2X\textsubscript{7} is the plasma membrane molecule that is responsible for the long known but little understood permeabilization of the plasma membrane consequent to stimulation of many cell types with extracellular ATP (Rozengurt et al., 1977; Cockcroft and Gomperts, 1979; Steinberg et al., 1987; Di Virgilio and Steinberg, 1993), the physiological function of this process has remained unknown.

It has been previously suggested that a possible role of P2Z/P2X\textsubscript{7} receptor could be in cellular communication, in a gap junction-like fashion (Steinberg et al., 1990; Di Virgilio et al., 1995). Two years ago we provided preliminary evidence in support of this hypothesis by observing that specific blockade of this receptor with oxidized ATP (oATP) almost completely inhibits formation of multinucleated giant cells (MGCs),\textsuperscript{1} triggered by incubation of human macrophages with Con A and interferon-\gamma (IFN-\gamma; Falzoni et al., 1995). Treatment with oATP on the other hand did not affect chemotaxis, cell aggregation, or expression of the adhesion molecules CD11a, CD18, and CD54.

To further investigate the role of P2Z/P2X\textsubscript{7} receptor, we have selected J774 macrophage cell clones that express this

\textsuperscript{1} Abbreviations used in this paper: IFN-\gamma, interferon-\gamma; MGC, multinucleated giant cells.
receptor at different levels, from virtually none (P2Zhypocells) to very high levels (P2Zhyper cells). Experiments reported in this paper show that P2Zhyper macrophages become exceedingly fragile and susceptible to ATP-mediated cell death and spontaneously fuse during in vitro culture.

Materials and Methods

Cells

J74 mouse macrophages and P2Zhyper and P2Zhypocells were grown in DME supplemented with 10% heat-inactivated horse serum, penicillin (100 U/ml), and streptomycin (100 μg/ml) (complete DME medium). P2Zhypocells were selected by repeated rounds of incubation in the presence of 5 mM ATP, followed by cloning by limiting dilution. P2Zhyper variants were obtained by cloning by limiting dilution and selection of the clones that showed a higher ATP-dependent uptake of lucifer yellow. Stable transfectants of HEK293 cells expressing the rat P2X<sub>7</sub> or P2X<sub>5</sub> receptors were described previously (Evans et al., 1995; Surprenant et al., 1996) and were grown in DME F12 medium supplemented with 10% FCS and 300 μg/ml of G418 (Inalco, Milan, Italy).

Phase Contrast and Fluorescence Microscopy

Phase contrast and fluorescence photographs were taken with an inverted fluorescence microscope (Olympus IMT-2; Olympus Optical Co., Ltd., Tokyo, Japan) equipped with 40 and 100× (oil immersion) objectives and fluorescein and rhodamine filters.

Transmission Electron Microscopy

Cell monolayers were fixed in 2.5% glutaraldehyde in 0.1 M cacodylate buffer (pH 7.2–7.4) and post fixed in 1% OsO<sub>4</sub> in the same buffer. Samples were then dehydrated and embedded in Araldite Durcupan (Fluka Chemie AG, Buchs, Switzerland). Blocks were cut with a microtome (Ultracut S; Reichert, Vienna, Austria), and ultra-thin sections were stained with uranyl acetate and lead citrate with an Ultrastainer (Reichert). Examination was performed in a transmission electron microscope (H-800; Hitachi Instr., San Jose, CA).

In Situ Hybridization

J74 mouse macrophages, P2Zhyper, and P2Zhypocells were allowed to adhere to polylysine-coated coverslides for 2 h. Preparations were fixed with 3% paraformaldehyde and incubated with either sense or anti-sense digoxigenin-UTP-labeled cRNA probes at a probe concentration of 50 ng/ml (Schaeren-Wiemers and Gerfin-Moser, 1993). After 16 h at 72°C, hybridization was detected with an anti-digoxigenin antibody conjugated to alkaline phosphatase (Boehringer Mannheim, Indianapolis IN) using 4-nitroblue tetrazolium chloride and 5-bromo-4-chloro-3-indolylphosphate (Boehringer Mannheim). Results are shown for an 18-h phosphatase reaction. The P2X<sub>7</sub> riboprobe corresponds to nucleotides 1,150–1,900 of the corresponding cDNA.

Western Blotting

Cells were detached from the tissue culture using a scraper and were suspended in PBS in the presence of protease inhibitors (phenylmethylsulfonyl fluoride, 4 mM; pepstatin, 2 mg/ml; leupeptin, 2 mg/ml; trypsin inhibitor, 2 mg/ml; aprotinin, 2 mg/ml). Cells were freeze thawed three times, alternating dry ice and 37°C water bath, and membranes were pelleted. Proteins were extracted in lysis buffer (10 mM Tris-HCl, pH 8.5, 150 mM NaCl, 1 mM CaCl<sub>2</sub>, 2% Triton X-100) in the presence of protease inhibitors. The extracts were centrifuged at 40,000 rpm to remove insoluble material. Equal concentrations of protein extracts were separated on an 8% polyacrylamide gel in the presence of SDS and transferred to nitrocellulose membrane.

Nitrocellulose membranes were incubated with primary antibody followed by peroxidase-conjugated swine anti-rabbit immunoglobulins (DAKO Spa, Milan, Italy) and developed using the chemiluminescent substrate ECL (Amersham Intl., Little Chalfont, UK).

Antibody

The rabbit polyclonal anti-P2X<sub>7</sub> antiserum was raised against the synthetic peptide corresponding to the last 20 amino acids of the rat P2X<sub>7</sub> protein (Collo, G., S. Neidhart, E. Kawashima, M. Kosco, R.A. North, and G. Buell, manuscript submitted for publication).

Results

Phenotypic Characterization of P2Zhyper and P2Zhypocells

P2Z/P2X<sub>7</sub> receptor function can be demonstrated by the ATP-mediated uptake of extracellular hydrophilic fluorescent markers such as lucifer yellow or YO-PRO. Fig. 1 shows that incubation in the presence of 3 mM ATP triggered a massive uptake of lucifer yellow by P2Zhyper cells (Fig. 1, A and D), while on the contrary almost no uptake by P2Zhypocells was detectable (Fig. 1, C and F). P2Zwt showed an intermediate behavior (Fig. 1, B and E). Nearly all P2Zhyper cells were positive for lucifer yellow uptake to varying levels. Because the sustained activation of the P2Z/P2X<sub>7</sub> receptor leads to cell death, we tested for the release of the cytoplasmic enzyme lactic dehydrogenase in response to ATP. Whereas P2Zhypocells were unaffected, P2Zhyper rapidly released lactic dehydrogenase, and P2Zwt again showed an intermediate behavior (Fig. 2).

P2Zhyper cells overexpress the P2Z/P2X<sub>7</sub> mRNA and protein as seen by in situ hybridization (Fig. 3) and western (Fig. 4) blot analysis. P2Zhyper macrophages (Fig. 3, A) show a very strong reactivity to the specific P2X<sub>7</sub> anti-sense riboprobe, while P2Zhypocells (Fig. 3, B) have little, if any, reactivity with the notable exception of one highly reactive cell. J774 wild-type cells (Fig. 3, C) show an intermediate reactivity. A stable transfectant (HEK293 cells) for rat P2X<sub>7</sub> is shown for comparison (Fig. 3, D). On average, we found that in the P2Zhyper population ~83 ± 5% of the cells were strongly positive, as compared to ~65 ± 8% of positive cells in the J774 wild-type population and only 2–3% of positive cells in the P2Zhypocells. No hybridization was detected with the sense riboprobe (not shown). Fig. 4 shows that the specific polyclonal Ab raised against the carboxy-terminal tail of the receptor recognized in J774 wild-type cells a protein of the expected molecular weight of ~72. This band was stronger in P2Zhyper cells and absent in P2Zhypocells and control cells.

Spontaneous Cell Fusion in P2Zhyper Monolayers

P2Zhyper cells were very fragile and easily died upon reaching confluence. Nonetheless, in many cultures we observed that shortly after reaching confluence (usually 3 d after plating), macrophages formed dense aggregates in which MGCs were easily detected (Fig. 5). During the last two years we have selected >20 P2Zhyper cell clones, and invariably, although to a different extent, we have observed formation of MGCs in all these cell populations. MGCs had different shapes: round, with short pseudopods (Fig. 5 A); star-like, with dendritic-like elongations (Fig. 5 B); polygonal, with a regular cellular contour (Fig. 5 C). MGCs contained from 2 to >10 large nuclei usually lo-
cated in the center and were also readily observed in monolayers of HEK293 cells stably transfected with the P2X7 receptor cDNA (Fig. 5D); on the contrary, fused cells were never observed in monolayers of P2Zhypo or P2Zwt macrophages (Fig. 6). In principle, MGCs could originate either from a real membrane fusion event or from kariokinesis without cell division. To solve this issue, we pooled two batches of P2Zhyper cells that had been previously labeled with different endosome–lysosome markers, lucifer yellow, and Texas red. Real fusion events were detected by observing both yellow/green and red vesicles within the same MGC. Two examples of such fusion are shown by arrowheads, which identify MGCs of different shapes in Fig. 7. On the contrary, individual macrophages are labeled by either the yellow/green or red tracer (Fig. 7, closed triangles). Obviously, cell fusion can also occur between macrophages labeled with the same tracer, such as the spindle-shaped MGC visible in the left-hand side of Fig. 7A (closed diamond), that is stained only by Texas red.

P2Zhyper macrophages formed cell aggregates that preceded fusion. It was very common to find within such aggregates, images suggestive of initial cytoplasmic communication between adjacent cells (Fig. 8, A and B, arrows).
Most often, an already-formed MGC was itself a center of aggregation to which other cells were attracted up to the eventual fusion (Fig. 8 C). We examined some of these aggregates by transmission electron microscopy. As shown in Fig. 9, there were sites of close apposition between membranes of adjacent cells, and sometimes the juxtaposed plasma membranes were about to fuse (Fig. 9, arrows).

Our group introduced a few years ago the only available inhibitor of P2Z/P2X<sub>7</sub> receptor, the dialdehyde compound oATP (Murgia et al., 1993). This reagent covalently binds to the receptor and causes an irreversible inactivation. Pretreatment of P2Z<sub>hyper</sub> cells with oATP for 2 h completely blocked spontaneous MGC formation without inhibiting aggregation (Fig. 10). Similar inhibition has been reported for monocyte-derived human macrophages (Falzoni et al., 1995). It has been recently suggested (Baricordi et al., 1996; Buell et al., 1996) that during in vitro culture, immune cell P2X receptors may be in a state of chronic desensitization due to a continuous leakage of ATP into the extracellular milieu. This hypothesis has received strong support from the recent demonstration that macrophages and microglial cells can release ATP via a nonlytic mechanism (Ferrari et al., 1997). Therefore, we tested the effect of the ATP-consuming enzyme hexokinase on MGC formation in P2Z<sub>hyper</sub> monolayers. Treatment with 100 μg/ml of hexokinase almost doubled the number of MGCs, from 6±3 to 12±2/field, as quantitated by observing 4 randomly selected microscopic fields from 2 different wells for each experimental condition, i.e., control and hexokinase-treated monolayers.

An interesting question is the fate of MGCs. Within 24–36 h after formation, MGCs started to deteriorate: cytoplasmic vacuoles appeared, nuclei condensed, and finally the cell body fragmented (Fig. 11, A–D). An MGC’s fate did not depend on the age of the culture but rather on that of the individual MGC; no MGC survived longer than 72 h after fusion.

**Discussion**

It is increasingly appreciated that ATP is an important mediator of intercellular communication (Burnstock, 1972, 1996; Zimmermann, 1994; Di Virgilio et al., 1996). Biological effects of ATP as an extracellular messenger are mediated through P2 purinergic receptors. Macrophages express both G protein–coupled P2Y and intrinsic ion channel P2X receptors (Greenberg et al., 1988). An intriguing P2X receptor expressed by macrophages is P2Z/P2X<sub>7</sub>, whose most characteristic property is to form an aqueous pore that, depending on the ATP dose and frequency of stimulation, forms an ion channel or a non-
selective pore that admits hydrophylic molecules of molecular mass up to 900 D (Steinberg et al., 1987; Di Virgilio et al., 1995). This receptor is specifically activated by ATP, probably in its ATP4\(^{2-}\) form, like the ATP-gated pore of mast cells (Cockcroft and Gomperts, 1979), and inhibited by the dialdehyde derivative oATP (Murgia et al., 1993). The only selective agonist, the ATP derivative benzoylbenzoic ATP, is about one order of magnitude more potent than ATP (Nuttle and Dubyak, 1994).

The only well characterized and generally accepted effect consequent to P2Z/P2X\(_7\) stimulation is cell death (Murgia et al., 1992; Molloy et al., 1994; Blanchard et al., 1995). Opening of the P2Z/P2X\(_7\) pore causes large transmembrane ion fluxes leading to Na\(^+\) and Ca\(^{2+}\) overflooding and K\(^+\) depletion accompanied by loss of cytoplasmic low molecular weight metabolites (Steinberg and Silverstein, 1987). Short permeabilizations (up to 10–15 min) are in general harmless, in that once extracellular ATP is removed, the pore reseals, and intracellular ion homeostasis is re-established. However, a sustained activation is invariably followed by cell death by either necrosis or apoptosis, depending on the length of the stimulation and the ATP concentration (Murgia et al., 1992; Molloy et al., 1994).

Why macrophages and a few other cell types should express such a dangerous receptor is not clear. It might be assumed that the receptor is never active, because in the body the extracellular ATP concentrations necessary for its opening (hundreds of micromolar or millimolar) are never reached. However, this is clearly not the case (Hoyle and Burnstock, 1996). More and more cell types are shown to be capable of releasing ATP via a number of different mechanisms. The most straightforward is exocytotic release of granular ATP by platelets, adrenal chromaffine cells, or neurons (Born and Kratzer, 1984; Zimmermann, 1994), but nonexocytotic mechanisms have also been im-

**Figure 5.** Spontaneous formation of multinucleated giant cells in monolayers of P2Zhyper cells and HEK293 cells transfected with the P2X\(_7\) receptor. Cells were plated at the concentration of \(5 \times 10^5\) cells/ml and grown for 3 d. (A–C) P2Zhyper macrophages; (D) HEK293 cells stably transfected with the P2X\(_7\) receptor. Photographs were taken with a 20\(\times\) objective in A and a 40\(\times\) objective in B-D. Bars: (A) 50 \(\mu\)m; (B–D) 25 \(\mu\)m.

**Figure 6.** Lack of formation of multinucleated giant cells in monolayers of P2Zwt and P2Zhypo macrophages. P2Zhypo (A) and P2Zwt (B) macrophages were plated as described in Fig. 5 and photographed with a 20\(\times\) objective. Bar, 50 \(\mu\)m.
 complicated (Pearson and Gordon, 1979; Gordon, 1990). For example, some membrane transporters such as P-glycoprotein and the CFTR protein have been proposed to carry ATP as well as other smaller ions (Abraham et al., 1993; Reisin et al., 1994; Schwiebert et al., 1995; but see Reddy et al., 1996, for contrasting results). In addition, because cytoplasmic ATP concentration is in the 5–10 mM range, any damage of the plasma membrane is likely to cause substantial release of ATP in the extracellular milieu. Events able to release ATP can range from mild injuries such as shear stress applied to the vessel wall to outright necrotic lesions. Under normal physiological conditions
these events could be uncommon, but at sites of acute or chronic inflammatory reactions they might be frequent and thus contribute significantly to the accumulation of high concentrations of extracellular ATP.

Our lab and others originally proposed that the P2Z/P2X7 receptor could have a role as a suicide receptor exploited by the immune system to eliminate unwanted cells and more generally down regulate the immune response (Di Virgilio et al., 1989, 1990; Filippini et al., 1990). This hypothesis is clearly very speculative, but a number of groups have already provided data in its support (Blan-
A cytokine that is massively released during the immune and inflammatory responses. Under these conditions, macrophages are known to establish close contact with each other and with different cell types, such as helper or cytotoxic lymphocytes, and to release several cytokines. Thus, it can be hypothesized that the P2Z/P2X7 receptor could be involved in intercellular communication, maybe to help establish bridges between the cytoplasm of adjacent macrophages, in a gap junction-like fashion. This process might contribute to formation of MGCs, a typical histologic feature of granulomatous inflammation. MGCs are known to arise from the fusion of adjacent macrophages and not from endomitosis (i.e., nuclear division without cytoplasmic division; McInnes and Rennick, 1988; Takashima et al., 1993), but the mechanism and the membrane molecules involved are still unknown. Several cytokines (e.g., IFN-γ, TNF-α, IL-3, and IL-4) have been implicated in different stages of MGC formation, but no clear conclusion has been reached about the identity of the fusogenic cytokine (Weinberg et al., 1984; Most et al., 1990). Molecules of the LFA-1 family might be involved, but again no firm evidence has been provided (Most et al., 1990).

J774 macrophage cells have been instrumental in the identification and characterization of the P2Z/P2X7 receptor. Steinberg and Silverstein performed a thorough investigation of the pharmacological and physiological properties of this molecule and introduced the ATP-resistant clones that became an invaluable tool at a time in which the cDNA was not available (Steinberg and Silverstein, 1987; Steinberg et al., 1987; Greenberg et al., 1988). We have in our laboratory ATP-resistant clones that were selected 10 yr ago. The cellular basis for the ability to select stable ATP-resistant cell clones rests on the marked heterogeneity of the wild-type J774 population with respect to P2Z/P2X7 expression. In general, susceptibility to ATP-mediated permeabilization of J774 cells assessed by uptake of hydrophobic fluorescent markers ranges from zero (dark cells) to very high (bright cells), with the full range of intermediate conditions. Very bright J774 cells can be selected, by limiting dilution, and clonally expanded. As previously shown by our group (Chiozzi et al., 1996), J774 hyper clones thus selected not only undergo a massive ATP-dependent lucifer yellow uptake but also cell death, confirming that high expression of the P2Z/P2X7 receptor confers an unusual sensitivity to ATP cytotoxic effects. P2Zhyper macrophages turned out to be very delicate cells to grow in culture, very probably as a direct consequence of high expression of the P2Z/P2X7 receptor, but very interestingly they also underwent a high rate of spontaneous fusion after 2 or 3 d of in vitro culture. Al-

Figure 11. Multinucleated giant cells die after fusion. P2Zhyper macrophages were plated as described in Fig. 5. Sequential pictures were taken from the same field at ~12-h interval.
though fusion was at its peak 3 d after plating, scattered MGCs could be observed even 1 d after seeding. Furthermore, HEK293 cells transfected with P2X7 cDNA, but not the wild-type control population, also showed several MGCs formation. These observations suggest that the mere hyperexpression of P2Z/P2X7 receptor may be sufficient to drive fusion in the absence of secreted or diffusible factors. High rate of spontaneous fusion correlated with high expression of P2Z/P2X; mRNA and reactivity with specific Abs; furthermore, fusion was abolished by αATP, the only selective inhibitor so far available. These observations, together with earlier convergent reports from our laboratory (Falzioni et al., 1995), point to a crucial role of this receptor in macrophage fusion, although they do not allow us to pinpoint exactly at what stage of the fusion process the receptor is involved.

Once formed, MGCs have a very short life. Within a few hours the cytoplasm becomes filled with phase-lucent vacuoles, fuzzy, and fragmented. At the end of this process, pinpoint exactly at what stage of the fusion process the re-

This report provides further evidence in support of the hypothesis that the pore-forming P2Z/P2X7 receptor participates in a crucial step of macrophage fusion leading to MGCs formation, a role of potential interest in the evolution of chronic granulomatous inflammation.

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