Application of ADA1 as a new marker enzyme in sandwich ELISA to study the effect of adenosine on activated monocytes

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Enzyme-linked immunosorbent assay (ELISA) is a valuable technique to detect antigens in biological fluids. Horse radish peroxidase (HRP) is one of the most common enzymes used for signal amplification in ELISA. Despite new advances in technology, such as a large-scale production of recombinant enzymes and availability of new detection systems, limited research is devoted to finding alternative enzymes and their substrates to amplify the ELISA signals. Here, HRP-avidin was substituted with the human adenosine deaminase (hADA1)-streptavidin complex and adenosine as a detection system in commercial ELISA kits. The hADA1 ELISA was successfully used to demonstrate that adenosine, bound to A1 and A3 adenosine receptors, increases cytokine secretion by LPS activated monocytes. We show that hADA1-based ELISA has the same sensitivity, and also provides identical results, as HRP ELISA. In addition, the sensitivity of hADA1-based ELISA could be easily adjusted by changing the adenosine concentration and the incubation time. Therefore, hADA1 could be used as a detection enzyme with any commercial ELISA kit with a wide range of concentration of antigens.

Enzyme-linked immunosorbent assay (ELISA) is a very popular test that uses antibodies and the enzyme-mediated reaction to detect the antigen of interest in a fluid sample1–3. Among various types of ELISAs, a sandwich ELISA is highly specific, recognizing two different epitopes on the same antigen with two types of antibodies, i.e. a capture antibody attached to a solid phase, such as 96-well microtiter plates, and a detection antibody that is biotinylated4. In order to detect constituents present in low amounts, it is often necessary to devise procedures capable of strongly amplifying the enzymatic signal. Horseradish peroxidase (HRP) and alkaline phosphatase (AP), which were introduced in the 1970s, are the most commonly used enzymes to bind the detection antibody for a signal amplification5,6. Multiple chromogenic and fluorescent substrates were designed to detect the activity of HRP and AP in ELISA kits4. However, many of them are quite expensive, unstable, and potentially harmful. The last few decades have brought in new technology, both in large scale expression of recombinant enzymes and development of new detection systems, such as the UV spectrum plate readers. Adenosine deaminase (ADA, EC 3.5.4.4) is a key enzyme of the purine metabolism, catalyzing the irreversible deamination of adenosine or deoxyadenosine to inosine or deoxyinosine, respectively. ADA has been found in bacteria, plants, and animals7. In humans, there are two ADAs, hADA1 and hADA28. Recently, we have developed a simple assay to detect hADA2 concentrations in biological fluids9. According to this assay, hADA2 is captured with polyclonal antibodies, which are absorbed by a 96-well microtiter plate, and then adenosine is added to detect ADA2 enzymatic activity. The amount of inosine produced is further analyzed on a separate detection plate, and can be followed by an increase in the absorbance ratio at 245 and 265 nm. The ratio is proportional to the hADA2 concentration, and hence, the concentration of ADA2 in a sample can be determined using hADA2 standards. In contrast to 120 kDa hADA2, hADA1 is a small 40 kDa enzyme, with higher turnover rate (kcat) and lower Km for adenosine8,10. Therefore, hADA1 and adenosine could be potentially used in the ELISA amplification system, applying a similar detection assay designed for hADA2.

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To validate ADA1 ELISA, we studied the effect of low concentrations of adenosine on the cytokine secretion by monocytes in response to their activation with Lipopolysaccharides (LPS). It is known that the activation of human monocytes with LPS induces expression of A1, A2A, A2B, and A3 adenosine receptors. According to the literature, adenosine is thought to be largely an immunosuppressive molecule. It was shown that adenosine at a high concentration binds to A2A receptors expressed on activated monocytes, and decreases the level of cytokines secretion, such as TNF-α. However, little is known about the role of A1 and A3 receptors in the regulation of monocytes activity, and the effect of low adenosine concentration on the cells’ activation. Here, using selective and non-selective adenosine receptor antagonists, we investigated the effect of low-level adenosine in stimulation of cytokine secretion by LPS-activated monocytes. To measure the cytokine secretion, we used a standard ELISA with HRP or hADA1 as detection enzymes, and showed that hADA1 could substitute HRP in commercial ELISA kits.

**Results**

**ADA1 expression and purification.** Wild-type hADA1 was expressed in the cytoplasm of HEK 293T cells stably transfected with lentiviral particles, as described in Methods (Fig. 1A). The enzyme was purified with a high yield from the cell lysate using a simple two-step chromatography (Fig. 1D). SDS-PAGE analysis confirmed the high purity of the enzyme (Fig. 1B). Enzymatic analysis showed that hADA1 had similar catalytic parameters.
Recombinant hADA1 was further chemically biotinylated without any loss of its enzymatic activity, indicating that the biotinylated hADA1 might be used to amplify the signals in standard ELISA.

**Design and sensitivity of ADA1 ELISA.** In the next series of experiments, we set up hADA1 ELISA using commercially available kits from BioLegend, to prove that hADA1 can be used as a detection enzyme. The antigen, bound to monoclonal capture antibodies, which were attached to the surface of 96-well microtiter plates, was further linked to biotinylated detection antibodies following a standard protocol of the ELISA kit (Fig. 2A). In the next step, hADA1-streptavidin complex was added to the plate. Subsequently, after the incubation with hADA1-streptavidin, adenosine was added to start the reaction of adenosine deamination by hADA1. The reaction was stopped by a transfer of an aliquot of the reaction mix to a separate detection plate, and the concentration of inosine was detected using three different assays. The first, so-called UV assay, was originally developed to measure the concentration of hADA2 in human plasma. Accordingly, the reaction mix from the first plate was transferred to a UV transparent 96-well plate, containing water to dilute the nucleosides to 0.1 mM concentration, and then the UV-transparent plate was read at 245 and 265 nm on a Synergy H1 hybrid Reader (Biotek). The reaction of deamination was followed by an increase in the ratio between the absorbances at 245 and 265 nm.

$$k_{cat} = 190 \text{ s}^{-1} \text{ and } K_m = 26 \mu \text{M}$$

(10) to the recombinant hADA1 expressed in bacteria (Fig. 1C). Recombinant hADA1 was further chemically biotinylated without any loss of its enzymatic activity, indicating that the biotinylated hADA1 might be used to amplify the signals in standard ELISA.

Figure 2. Design of hADA1 ELISA. (A) A schematic representation of the hADA1-based ELISA; (B,C) Standard curves for TNF-α, obtained using hADA1-streptavidin in standard ELISA. The concentration of TNF-α is proportional to the concentration of adenosine converted to inosine. (B) The reaction of adenosine hydrolysis to inosine by hADA1 is monitored by the absorbance ratio between 245 and 265 nm. (C) The amount of inosine produced is detected in the enzymatic color reaction, resulting in the increase in the absorbance at 550 nm. Each dot represents the average of two replicates.
and 265 nm. The ratio was proportional to the concentration of hADA1 and antigen (Fig. 2B). This simple assay doesn't require any additional reagents, and the reaction mix from the first plate can be analyzed multiple times (Fig. 3). Alternatively, in a color assay, the reaction of adenosine deamination could be monitored by a method wherein inosine is subsequently converted by purine nucleoside phosphorylase (PNP) and xanthine oxidase (XOD) to uric acid and hydrogen peroxide, which is further reacted with N-Ethyl-N-(2-hydroxy-3-sulfopropyl)-3-methylaniline (EHSPT) and 4-aminoantipyrine (4-AA) in the presence of peroxidase (POD) to generate quinone dye. Quinone dye was detected at 550 nm in the visible spectrum, which allows using standard 96-well plates and a simple plate reader (Fig. 2C). In the third assay, the sensitivity of inosine detection could be significantly increased by substituting EHSPT and 4-AA with a fluorescent reagent, Ampliflu Red. Accordingly, Ampliflu red is converted into a highly fluorescent compound, resorufin, and the reaction of adenosine deamination is monitored by the increase in fluorescent intensity (Fig. 4B). The sensitivity of the hADA1 assay for a given ELISA kit or antigen concentration can be adjusted either by changing adenosine concentration or incubation times (Fig. 3). In general, the increase in the incubation time and the use of lower adenosine concentrations allow reaching the maximum sensitivity of the assay. Therefore, hADA1 ELISA could be used for a wide range of antigen concentration, ranging from nanograms to picograms per milliliter (Fig. 4A). The use of fluorescent reagents adds even higher sensitivity to the assay (Fig. 4B). Biotinylated human ADA1 is very stable and it could be kept at +4°C in a phosphate-saline buffer containing sodium azide. The enzyme remains fully active for an extended period of time providing reliable and reproducible results (Fig. 5).

Study of monocytes activation in the presence of ADORs antagonists using a sandwich ELISA with either HRP or hADA1. It has been shown that monocytes activation is controlled by adenosine. Here, we used LPS-activated monocytes to study the effect of different ADOR antagonists on the expression level of TNFα or MCP-1. To compare the hADA1-based ELISA system with a standard HRP ELISA, the concentration of cytokines in the cell supernatants were determined in parallel, using either HRP or hADA1 as a detection enzyme. As shown in Fig. 6A, the results obtained with HRP and hADA1 were almost identical. The addition of a non-specific adenosine receptor antagonist to the LPS-activated monocytes leads to a decrease in both TNF-α and MCP-1 secretion by the cells. This indicates that, in the absence of ADOR antagonist, adenosine, which is present in a test tube, binds to adenosine receptors and stimulates secretion of the cytokines by activated cells. Moreover, the release of MCP-1 by the cells is completely abolished when adenosine receptors are inactivated by

Figure 3. Sensitivity of hADA1 ELISA can be controlled by adenosine concentration and incubation time. (A,B) Standard curves obtained for TNF-α and MCP-1 ELISA at different concentrations of adenosine after 20 hours of incubation. (C,D) Standard curves obtained for TNF-α and MCP-1 ELISA at different times of incubation with 800 μM of adenosine after 20 hours of incubation. Each dot represents the average of two replicates.
ADOR antagonists. It is known that, at low adenosine concentration, only A1 and A3 receptors can bind adenosine\(^1\). The binding to A2A receptor requires higher concentrations of adenosine, because this receptor has a lower affinity for the ligand. Thus, we used specific A1, A3, and A2A ADOR antagonists to reveal the receptors that are involved in the regulation of monocytes activation. As seen in Fig. 6, the addition of A2A antagonist didn’t affect the level of both cytokines secretion, suggesting that this receptor is not involved in the cells’ activation at low levels of adenosine. Conversely, A3 antagonist almost completely inhibited MCP-1 secretion by LPS-activated monocytes. There was a negative effect on TNF-\(\alpha\) secretion as well. The effect of the A1 antagonist on MCP-1 release from the activated monocytes was negligible, while the level of TNF-\(\alpha\) secretion was significantly diminished. These results show that, at low concentration, adenosine binds to A3 and A1 receptors, and stimulates the cytokines secretion by activated monocytes.

**Discussion**

To date, a few enzymes have been employed as markers in the enzyme immunoassays\(^4\). Horseradish peroxidase, alkaline phosphatase, and \(\beta\)-galactosidase were the main enzymes employed to amplify signals in the enzyme-linked detection assays. One of the reasons for the scarce choice of the enzymes is the high requirements placed on them, such as a high specific activity of the enzyme after the labeling, availability of the purified enzyme at low cost, high stability, availability of inexpensive and stable substrates, and their stable chromogenic and fluorogenic products\(^4\). Horseradish peroxidase is ideal in many respects for these applications, because it is smaller, more stable, and less expensive than other popular alternatives, such as alkaline phosphatase. It also has a high turnover rate that allows generation of strong signals in a relatively short time span. Therefore, the majority of commercial ELISA kits utilize HRP as a detection enzyme. There are a variety of HRP substrates available for chromogenic, chemiluminescent, and chemiluminescent imaging. However, a low stability, nonspecific color precipitation and toxicity could be the main issues for the selected substrates. Another inconvenience is that each company tends to produce their own HRP conjugates and detection substrates, which are not interchangeable between the kits of different companies. A short range of the antigen concentration that could be detected with...
a given ELISA kit often results in the loss of valuable samples, because the concentration of the antigen in the samples doesn't fit a standard curve. Here, we describe an alternative detection system, with human ADA1 as a detection enzyme, which has several advantages over HRP ELISA. The use of two separate plates for the antigen binding and the analysis of enzymatic reaction gives more flexibility, allowing the use of hADA1 ELISA kits with UV, fluorescent, and visible spectrum plates. It also allows achieving the desired sensitivity of the kit to get an accurate estimate of the antigen concentration in the sample.

At first, we developed a simple technology to produce large quantities of recombinant human ADA1 (Fig. 1D). The enzyme was expressed in 293T HEK cells transfected with lentiviral particles containing hADA1 gene (Fig. 1A). To the best of our knowledge, this is the first report on purification of fully active human recombinant ADA1 from mammalian cells. Human ADA1 is not toxic to the cells and, thus, it is accumulated at high concentrations in the cell cytoplasm. An easy three-step purification protocol permits isolating highly active and pure hADA1 (Fig. 1B). The chemical biotinylation of the enzyme didn't affect the enzyme activity, which was preserved for at least one year (Fig. 5). In contrast to HRP, which is inhibited by sodium azide, hADA1 could be stored at 4°C in PBS containing NaN3. Biotinylated hADA1 was simply coupled to streptavidin by mixing of two proteins. Since hADA1 is a small 40 kDa enzyme, the streptavidin-hADA1 complex can be bound to biotinylated detection antibodies without any steric hindrance (Fig. 2). The enzymatic reaction was started by adding a natural ADA substrate adenosine in PBS buffer. The main advantages of adenosine are that it is cheap, soluble, stable, and non-toxic. It could either be stored in solution or prepared freshly by dissolving in PBS. The reaction of adenosine deamination, catalyzed by hADA1, was stopped by transferring an aliquot of the reaction mix to a detection plate (Fig. 2A). Due to the difference in the absorption spectrum between adenosine and inosine, the

Figure 6. Effect of ADORs antagonists on TNF-α and MCP-1 secretion by LPS-activated monocytes. (A) Graphs showing the concentration of TNF-α and MCP-1 released by LPS-activated monocytes, which was determined either with a standard ELISA kit (HRP) or with hADA1 ELISA, using UV (Fig. 2B) and color (Fig. 2C) detection assays. The monocytes were incubated with 10 ng/ml LPS, either in the absence (test tube) or presence of nonselective A2A,A3A1A3 or selective A3, A2A and A1 ADOR antagonists. (B) A schematic representation of the experiment.
The advantage of the hADA1 based ELISA is that the reaction could be analyzed on a separate detection plate at different times of incubation with adenosine (Fig. 3C,D). This adds more flexibility over the HRP system (Table 1). Therefore, it is possible to adjust the sensitivity of the assay to get the highest accuracy for the antigen detection in the samples, by varying either adenosine concentration (Fig. 3A,B) or the incubation time with adenosine (Fig. 3C,D). This also allows adopting this system for use with any commercial ELISA kit. Moreover, the use of hADA1 instead of HRP permits the increase of the range of concentration of standards to detect the samples with a broad distribution of the antigen concentrations. For instance, the samples with high concentration of antigen could be read after a short incubation time, while the samples with low concentrations of antigens could be detected after a prolonged incubation with adenosine (Fig. 4A). Although it takes about 7 hours to reach a sensitivity comparable to HRP ELISA, the incubation time could be reduced to less than 1 hour by decreasing the concentration of adenosine 10 times. The detection diapason can be further extended by the use of Ampliflu Red fluorogenic reagent (Fig. 4B). Therefore, with the hADA1 detection system, there is no need to dilute samples to fit a narrow window of concentration, as required in the HRP-based ELISA kits. In addition, the completeness of the reaction of deamination can be checked by reading the standard curve together with a few key samples before the analysis of the whole plate. This procedure, which cannot be done with the HRP ELISA, increases the accuracy of the antigen detection and assures that the antigen concentration in all samples fits to the standard curve.

To validate hADA1 as a detection enzyme, we used both HRP and hADA1-based ELISA to measure the concentration of cytokines produced by LPS-activated monocytes in the cell culture medium. We studied the effect of low concentrations of adenosine, which accumulates in the cell culture, on the level of TNF-α and MCP-1 secretion by activated monocytes. To block the binding of adenosine to adenosine receptors (ADOR), we added either nonselective ADOR antagonist or selective A1, A2A or A2B antagonists. In the presence of nonselective ADOR antagonist, we saw a dramatic decrease in the release of both cytokines by activated monocytes (Fig. 6A). Furthermore, the level of MCP-1 was almost undetectable. This indicated that adenosine was present in our system, and that the blocking of adenosine receptors with ADOR antagonists prevented adenosine binding to ADORs. We used specific ADORs antagonists to find out which of the ADOR receptors bind adenosine and stimulate cytokine secretion by activated monocytes. It was shown that blocking of A2A receptor didn’t change the level of cytokines secretion into the culture medium, suggesting that the concentration of adenosine was too low to activate A2A receptor. However, the blocking of either A1 or A3 receptor decreased the concentration of cytokines released by activated monocytes. In addition, our results showed that TNF-α secretion was stimulated by adenosine binding to both A1 and A3 receptors, while the level of MCP-1 secretion was mainly controlled by A3. This suggests that the binding of adenosine at low concentrations to either A1 or A3 receptors stimulates the release of TNF-α and MCP-1 (Fig. 6B). This effect is opposed to that of high concentrations of adenosine, where secretion of TNF-α by LPS-activated cells is inhibited by adenosine bound to A2A receptors13. Therefore, a stimulatory effect of low concentrations of adenosine on the monocytes activation should be considered when ADOR antagonists are validated as potential drugs to tackle various immune disorders. In our experiment, both hADA1 and HRP ELISA gave identical results (Fig. 5), proving that ADA1-based ELISA could be used as a valuable tool to measure antigen concentrations in biological fluids.

In conclusion, in this work we demonstrate that ADA1 can be used as an alternative detection enzyme in commercial ELISA kits (Table 1). The main advantage of the ADA1-based ELISA is the use of a stable, cheap, and non-toxic substrate adenosine. The product of the enzymatic reaction, inosine, has a different absorbance spectrum than adenosine, allowing the enzymatic reaction to be followed by a change in a ratio between absorbance at 245 and 265 nm. The amount of adenosine produced can be detected on a separate detection plate at different

|                     | ADA                  | HRP                  |
|---------------------|----------------------|----------------------|
| Standard range for a single experiment | 10 pg/ml–5000 pg/ml * | 7.8 pg/ml–500 pg/ml * |
| Coefficient of variation (CV%)          | 2.1–6.5 *            | 4.8–11.7 *           |
| Composition and concentration of the substrate | 0.5–1 μM adenosine | TMB and H2O2 (proprietary information) * |
| Light sensitivity of the substrate       | no                   | yes                  |
| Long-term stability of the substrate     | yes                  | no                   |
| Can be used with buffers containing sodium azide | yes           | no                   |
| Reaction can be analyzed several times  | yes                  | no                   |
| Reaction can be repeated                 | yes                  | no                   |
| Reaction can be analyzed with a UV reader| yes                  | no                   |
| A chromogenic or fluorescent substrate must be present in the reaction mix | no           | yes                  |
| Interchangeability                      | Can be used with different commercial ELISA kit | Pre-optimized to work with the reagents in a particular kit |

Table 1. The main differences between ADA and HRP ELISA. *The parameters are given for MCP-1 ELISA kit (BioLegends). Coefficient of variation was calculated from the values of 4 different samples with 6 replicates of the same assay.
incubation time points with adenosine using ultraviolet spectroscopy, colorimetry, and fluorescent assays. This adds flexibility to the assay, resulting in a higher accuracy and a broader range of antigen concentration detected by commercial ELISA kits.

**Methods**

**Cloning and expression of the human ADA1.** All methods were carried out in accordance with the relevant institutional guidelines. To stably express human ADA1 in cells transfected with a replication-defective lentiviral vector, the open reading frames (ORFs) of the genes were PCR amplified with the following primers: (F) 5′-AGCTCGAGACCGTCCACCATGGCCCGAGACGCCCGCT-3′ and (R) 5′-ATGGATCCGTAGCTCGAGGTCTTGCCTGAC-3′ (hADA1). The PCR products were then subcloned into the pCR2.1-TOPO plasmid, excised by restriction digest with XhoI and BamHI, and ligated into a XhoIBamHI-digested self-inactivating (SIN) transfer plasmid (pHR-cPPT-hB7-SIN)19. HEK 293T cells were then transfected with the transfer, VSV-G envelope and transfection reagent20. Subsequently, lentiviral particles were purified by ultracentrifugation from the cell culture medium of these HEK-293T cells, and were used to infect new 293T cells. The recombinant human ADA1 was purified from the 293T cell lysates. The cells were grown in complete DMEM medium (Sigma-Aldrich), supplemented with 5% FBS, 100 U/ml penicillin, 100 μg/ml streptomycin, and 2 mM L-glutamine. The cells were trypsinized with trypsin/EDTA (Sigma-Aldrich), centrifuged for 5 min at 300 g, and the cell pellet was frozen in liquid nitrogen and kept at −80°C. Frozen 293T cells expressing hADA1 were lysed by three freeze/thaw cycles, using liquid nitrogen and a 42°C water bath. The cell lysate was re-suspended in 50 ml of ice-cold 50 mM Tris-HCl buffer (pH 6.8), 50 mM NaCl, 10 μM Zn(AcO)2, and 0.02% NaN3 (buffer A). The supernatant was separated from the cell debris by centrifugation at 4000 g for 20 min, filtered using 0.45 μm filters (Sigma-Aldrich), and applied onto a DEAE Sepharose (GE Healthcare) equilibrated with buffer A. The flow through containing ADA1 was collected, and the pH was adjusted to 8.4 with Tris base. The enzyme was applied onto a DEAE Sepharose column equilibrated with buffer B (50 mM Tris-HCl pH 8.5, 10 μM Zn(AcO)2, and 0.02% NaN3). hADA1 bound to the column was eluted using 0–500 mM NaCl gradient. The fractions containing ADA activity were pooled, concentrated using 10 kDa centrifuge ultraconcentrators (Millipore), and further purified on a Superdex 200 column (GE Healthcare), equilibrated with phosphate-buffered saline (PBS) containing 10 mM Zn(AcO)2 and 0.02% NaN3. The kinetic parameters of the recombinant human ADA1 and its activity were determined as described before8,9. Purified hADA1 was chemically biotinylated using a kit from G-Biosciences.

**Monocytes isolation and culture.** Human CD14+ monocytes were purified from buffy coats of healthy donors (Red Cross, Helsinki, Finland) by first isolating PBMCs with a Ficoll density gradient, and subsequently selecting for CD14+ monocytes using anti-CD14-conjugated magnetic microbeads (Miltenyi), as described previously (Zavialov et al., 2010a). The recovered cells were 95–99% CD14+, as determined by flow cytometry using the FITC conjugated anti-human CD14 mAb (BD Biosciences). RPMI 1640, supplemented with 1% non-essential amino acids, 1% sodium pyruvate, 100 U/ml penicillin, 100 μg/ml streptomycin, 2 mM L-glutamine, 0.1% 2-mercaptoethanol, and 10% FBS, was used as the complete medium in the cell cultures of monocytes. Monocytes were cultured in suspension in 5 ml polypropylene tubes (Falcon) at 0.5 × 106 cells/ml in complete medium of these HEK-293T cells, and were used to infect new 293T cells. The recombinant human ADA1 was purified from the 293T cell lysates. The cells were grown in complete DMEM medium (Sigma-Aldrich), supplemented with 5% FBS, 100 U/ml penicillin, 100 μg/ml streptomycin, and 2 mM L-glutamine. The cells were trypsinized with trypsin/EDTA (Sigma-Aldrich), centrifuged for 5 min at 300 g, and the cell pellet was frozen in liquid nitrogen and kept at −80°C. Frozen 293T cells expressing hADA1 were lysed by three freeze/thaw cycles, using liquid nitrogen and a 42°C water bath. The cell lysate was re-suspended in 50 ml of ice-cold 50 mM Tris-HCl buffer (pH 6.8), 50 mM NaCl, 10 μM Zn(AcO)2, and 0.02% NaN3 (buffer A). The supernatant was separated from the cell debris by centrifugation at 4000 g for 20 min, filtered using 0.45 μm filters (Sigma-Aldrich), and applied onto a DEAE Sepharose (GE Healthcare) equilibrated with buffer A. The flow through containing ADA1 was collected, and the pH was adjusted to 8.4 with Tris base. The enzyme was applied onto a DEAE Sepharose column equilibrated with buffer B (50 mM Tris-HCl pH 8.5, 10 μM Zn(AcO)2, and 0.02% NaN3). hADA1 bound to the column was eluted using 0–500 mM NaCl gradient. The fractions containing ADA activity were pooled, concentrated using 10 kDa centrifuge ultraconcentrators (Millipore), and further purified on a Superdex 200 column (GE Healthcare), equilibrated with phosphate-buffered saline (PBS) containing 10 mM Zn(AcO)2 and 0.02% NaN3. The kinetic parameters of the recombinant human ADA1 and its activity were determined as described before8,9. Purified hADA1 was chemically biotinylated using a kit from G-Biosciences.

**ELISA using hADA1.** The concentration of TNF-α or MCP-1 was determined using BioLegend ELISA kits according to the manufacturer’s instructions. For the experiments with ADA1-based ELISA, avidin-HRP was exchanged for hADA1-streptavidin complex following the protocol below:

1. Prepare 100 μg/ml hADA1-Streptavidin (based on the streptavidin concentration) by mixing hADA1 with Streptavidin in a molar ratio 0.5:1.
2. Dilute 100 μg/ml hADA1-Streptavidin to 500 ng/ml with 0.5% BSA, PBS, and 0.02% NaN3.
3. Add 100 μl of 500 ng/ml hADA1-Streptavidin to the wells of a 96-well plate, and incubate for 30 min at room temperature on a shaker.
4. Wash the plate 3 times with 200 μl 0.05% Tween 20 in PBS.
5. Wash the plate 1 time with 200 μl of PBS.
6. Add 100 μl of 800 μM adenosine in PBS, 5 μM Zn2+.
7. Incubate the plate at 37°C for 20 hours.
8. Transfer 25 μl of the reaction mix to a UV-transparent plate (Costar) containing 175 μl H2O2, to make up 200 μl of 0.1 mM adenosine. Read the plate on a Synergy H1 hybrid Reader (Biotek) at 245 and 265 nm to get the 265/245 ratio, and determine the concentrations from a standard curve. The UV-transparent plate can be reused multiple times.

Alternatively, 40 μl of the reaction mix were transferred onto a 96-well cell culture plate (Greiner) containing 60 μl of 0.0625 U/ml purine nucleoside phosphorylase, 0.025 xanthine oxidase, 0.01 U/ml horse radish peroxidase, 1.25 mM 4-aminoantipyrine, and 0.75 mM N-Ethyl-N-(2-hydroxy-3-sulphopropyl)-3-methylaniline (all from Sigma-Aldrich) in 0.1 M Tris-HCl pH 6.8. The plate was incubated at 37°C for 30 min, and the absorbance at 550 nm was determined using a Synergy H1 hybrid Reader (Biotek). For the detection using
a fluorescent reagent AmpliFlu Red (Sigma-Aldrich). 1.25 mM 4-aminoantipyrine and N-Ethyl-N-(2-hydroxy-3-sulfopropyl)-3-methylaniline were substituted with 0.1 mM AmpliFlu Red, and the fluorescent intensity (excitation 530 nm and emission at 590 nm) was determined using a Synergy H1 Hybrid Reader (Biotek).

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Author Contributions
A.Z. designed this work and analyzed the data. A.Z. and Y.L. supervised this study. A.Z., C.L., M.S. and C.W. performed the experiments. A.Z. prepared the manuscript. All authors read and approved the final manuscript.

Additional Information
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