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Abstract: Overexpression of the adenosine A1 receptor (A1AR) has been detected in various cancer cell lines. However, the role of A1AR in tumor development is still unclear. Thirteen A1AR mutations were identified in the Cancer Genome Atlas from cancer patient samples. We have investigated the pharmacology of the mutations located at the 7-transmembrane domain using a yeast system. Concentration–growth curves were obtained with the full agonist CPA and compared to the wild type hA1AR. H78L3.23 and S246T6.47 showed increased constitutive activity, while only the constitutive activity of S246T6.47 could be reduced to wild type levels by the inverse agonist DPCPX. Decreased constitutive activity was observed on five mutant receptors, among which A52V2.47 and W188C5.46 showed a diminished potency for CPA. Lastly, a complete loss of activation was observed in five mutant receptors. A selection of mutations was also investigated in a mammalian system, showing comparable effects on receptor activation as in the yeast system, except for residues pointing toward the membrane. Taken together, this study will enrich the view of the receptor structure and function of A1AR, enlightening the consequences of these mutations in cancer. Ultimately, this may provide an opportunity for precision medicine for cancer patients with pathological phenotypes involving these mutations.

Keywords: G protein-coupled receptors; adenosine A1 receptor; cancer; mutation; yeast system

1. Introduction

G protein-coupled receptors (GPCRs) are the largest protein superfamily in the human genome with approximately 800 subtypes [1]. They share structural characteristics of seven-transmembrane helices (TM) connected by an extracellular N-terminus, three extracellular loops (ELs), three intracellular loops (ILs), and an intracellular C-terminus [2]. GPCRs are widely distributed throughout the human body and regulate various crucial cellular and physiological functions by responding to a diverse set of endogenous ligands [3]. However, their aberrant activity and expression also substantially contribute to human pathophysiology [4].

Kinases, due to their central roles in the cell cycle, have been studied as a primary focus in preclinical oncology over the last two decades [5]. GPCRs, however, have been relatively under-investigated in this context, while an increasing amount of evidence shows that GPCRs act as regulators of tumor initiation and progression [6]. Malignant cells often hijack the normal physiological functions of GPCRs to survive, invade surrounding tissue, and evade the immune system [7]. Moreover, somatic mutations of GPCRs have been identified in approximately 20% of all cancers by a systematic analysis of cancer genomes [5].

The immune system plays a fundamental and essential role in the defense against cancer [8]. Adenosine, a nucleoside, and derivative of ATP, has emerged as a major
immune–metabolomic checkpoint in tumors [9]. Compared to healthy tissue, adenosine is accumulated over 50-fold in the hypoxic tumor environment, leading to a reduced anti-tumoral immune response [10]. Adenosine regulates various physiological effects and immune responses in cancer via adenosine receptors (ARs): the $A_1$, the $A_{2A}$, the $A_{2B}$, and the $A_3$ receptor [11]. Additionally, all ARs have been detected in different human tumor tissues [12]. Therefore, all four subtypes of ARs may regulate cancer progression in one way or another.

Growing evidence addresses the involvement of $A_1$AR in cancer progression, although its precise role is not well understood [13,14]. An increased expression level of the $A_1$AR has been detected in diverse cancer cells [15,16], where it appears to behave as both an anti- and pro-tumoral regulator in the development of different cancer types [10]. Interestingly, various single-site point mutations on $A_1$AR have been isolated from patients with different cancer types and collected by the TCGA Research Network (https://www.cancer.gov/tcga, 24 February 2022). Previous site-directed mutagenesis and docking studies on $A_1$AR have identified residues all over the protein involved in ligand recognition and/or functional activity [17,18]. Furthermore, several GPCR-conserved residues and motifs, for instance, the D2.50 residue, the ionic lock, the NPxxY motif, and the DRY motif, are located at 7-TM domains mediating ligand binding and signaling [19].

In this study, 13 mutations located at the 7-TM domains of the $A_1$AR have been selected from cancer patients using a bioinformatics approach. The effects of these mutant receptors on constitutive receptor activity and agonist-induced activation were tested in a ‘single-GPCR-one-G protein’ *S. cerevisiae* strain, which has been reported to be predictive of the mammalian situation [20,21]. Selected mutant receptors were further investigated for their effects on ligand binding and receptor activation in a mammalian system. Subsequently, we identified two CAMs, five CIMs, and six loss-of-function mutants (LFMs) based on the pharmacological effects of these mutant receptors. Thus, cancer-related mutations within the 7-TM domain may alter the role of $A_1$AR in cancer progression and the efficacy of drugs targeting $A_1$AR as a cancer therapeutic approach.

2. Results

2.1. Data Mining

Mutation data from cancer patient isolates were obtained by data mining the TCGA database on August 8, 2015. A total of 27 point somatic mutations were selected from (in total) 48 cancer-related point mutations of $hA_1$ARs based on selected cancer types, i.e., breast invasive carcinoma, colon adenocarcinoma, lung adenocarcinoma, lung squamous cell carcinoma, lymphoid neoplasm diffuse large B-cell lymphoma and rectum adenocarcinoma. After assigning Ballesteros Weinstein numbers to the positions by using the GPCRdb alignment tool, 13 mutations located at the 7-TM domains were selected for this study (Table 1). One mutation was located at the first, two at the second, two at the third, one at the fourth, one at the fifth, two at the sixth, and two at the seventh TM (Figure 1A).

2.2. Constitutive Activity of Mutant $hA_1$ARs

To first characterize the effect of the cancer-related mutations on the constitutive activity of the receptor, i.e., activity independent from an agonist, yeast growth assays were performed in the absence of the agonist. First, the optimal concentration of the histidine biosynthesis inhibitor (3-amino-1,2,4-triazole, 3-AT) for constitutive activity screening was determined in response to increasing concentrations of 3-AT (Figure 1B). Upon increasing concentrations of 3-AT, cell growth of both yeast cells transformed with a plasmid, with or without wild type $hA_1$AR, decreased (Figure 1B). At a concentration of 4 mM 3-AT, the two curves showed the largest differences in yeast growth; at this point, mutant receptors with increased constitutive activity (CAM) would show a higher growth level than wild type $hA_1$AR, while mutant receptors with decreased constitutive activity (CIM) would show a growth level in between wild type $hA_1$AR and empty vector. Thus, using this concentration of 3-AT provided the best window to screen for both CAMs and CIMs.
Table 1. List of cancer-related somatic mutations identified from different cancer types.

| Mutations | Cancer Types                   | Occurrence |
|-----------|--------------------------------|------------|
| A20T      | Colon adenocarcinoma           | 1          |
| A52V      | Breast invasive carcinoma      | 1          |
| D55V      | Breast invasive carcinoma      | 1          |
| D55G      | Colon adenocarcinoma           | 1          |
| H78L      | Lung adenocarcinoma            | 1          |
| P86L      | Rectum adenocarcinoma          | 1          |
| R122Q     | Colon adenocarcinoma           | 1          |
| L134F     | Lung squamous cell carcinoma   | 1          |
| W188C     | Colon adenocarcinoma           | 1          |
| S246T     | Breast invasive carcinoma      | 1          |
| T257P     | Lung adenocarcinoma            | 1          |
| S267I     | Colon adenocarcinoma           | 1          |
| G279S     | Colon adenocarcinoma           | 1          |

Cancer-related mutations showed various effects on the constitutive activity of the hA1AR (Figure 1C). Eleven out of the thirteen mutant receptors had a decreased constitutive activity compared to the wild type hA1AR. Among them, mutant receptors A52V, D55V, R122Q, L134F, W188C, and T257P even showed similar activities as yeast cells transformed by the empty vector. In contrast, increased constitutive activity was observed on two mutant receptors, i.e., H78L and S246T.

2.3. Agonist-Induced Receptor Activation of Mutant hA1ARs

To further characterize the activation profiles of these mutations, concentration–growth curves were determined in the presence of increasing concentrations of the selective hA1AR full agonist, CPA, and 7 mM 3-AT (Figure 2 and Table 2). Wild type hA1AR showed a
potency/pEC$_{50}$ value of $9.30 \pm 0.08$ and a maximum effect/E$_{\text{max}}$ value (ratio over wild type basal activity) of $4.83 \pm 0.30$ in the yeast system (Table 2).

![Figure 2](image)

Figure 2. Concentration–response curves of wild type and mutated hA$_1$ ARs. Data are separated for mutations located on (A) the first and second transmembrane helix, (B) third and fourth transmembrane helix, and (C) fifth, sixth, and seventh transmembrane helix. Data were normalized as ratio over basal activity of wild type hA$_1$AR (dotted line). Combined graphs are shown as mean ± SEM from at least three individual experiments performed in duplicate. Data for wild type are shown in dark blue, for CIMs in red, for CAMs in green, and for LFM in grey.

Table 2. Agonist (CPA)-induced receptor activation of wild type and mutant hA$_1$ARs in yeast liquid growth assays.

| Mutation | Basal | pEC$_{50}$ (−log M) | E$_{\text{max}}$ | Type |
|----------|-------|---------------------|------------------|------|
| Wild type | 1.00 ± 0.08 | 9.30 ± 0.08 | 4.83 ± 0.30 | -    |
| A20T$^{1.43}$ | 0.68 ± 0.14 | 9.24 ± 0.08 | 4.23 ± 0.60 | CIM  |
| A52V$^{2.47}$ | 0.24 ± 0.02 | 6.68 ± 0.09 | 1.86 ± 0.14 | CIM  |
| D55V$^{2.50}$ | 0.24 ± 0.04 | ND | ND | LFM  |
| D55C$^{2.50}$ | 0.50 ± 0.06 | ND | ND | LFM  |
| H78L$^{3.23}$ | 4.48 ± 0.12 | ND | 4.15 ± 0.17 | CAM  |
| P86L$^{3.31}$ | 0.28 ± 0.03 | ND | ND | LFM  |
| R122Q$^{4.40}$ | 0.57 ± 0.22 | 9.04 ± 0.14 | 4.67 ± 0.22 | CIM  |
| L134F$^{4.52}$ | 0.29 ± 0.04 | ND | ND | LFM  |
| W188C$^{5.46}$ | 0.32 ± 0.02 | 8.21 ± 0.10 | 4.35 ± 0.10 | CIM  |
| S246T$^{6.47}$ | 1.95 ± 0.27 | 9.42 ± 0.33 | 4.81 ± 0.26 | CAM  |
| T257P$^{6.58}$ | 0.24 ± 0.01 | ND | ND | LFM  |
| S267I$^{7.32}$ | 0.28 ± 0.01 | ND | ND | LFM  |
| G279S$^{7.44}$ | 0.33 ± 0.12 | 9.27 ± 0.09 | 4.96 ± 0.38 | CIM  |

Mutations are indicated using the numbering of the hA1AR amino acid sequence as well according to the Ballesteros and Weinstein GPCR numbering system [22]. All values are shown as mean ± SEM obtained from at least three individual experiments performed in duplicate. * Values were calculated as ratio over basal activity of wild type hA1AR. b Typing of the mutants was done according to their constitutive (in)activity and agonist-induced receptor activation. * p < 0.05; ** p < 0.01; *** p < 0.001; **** p < 0.0001 compared to wild type hA1AR, determined by a two-tailed unpaired Student’s t-test. ND: not detectable, CAM: constitutively active mutant, CIM: constitutively inactive mutant, LFM: loss of function mutant.

Almost half of the mutant receptors with decreased constitutive activity could not be activated by CPA anymore, namely D55V$^{2.50}$, D55C$^{2.50}$, P86L$^{3.31}$, L134F$^{4.52}$, T257P$^{6.58}$, and S246T$^{6.47}$, which resulted in typing them as loss of function mutants (Figure 2 and Table 2). Other mutant receptors with decreased constitutive activity could still be activated by CPA with equal or lower potency and efficacy values. Specifically, in response to CPA, mutant receptors A20T$^{1.43}$, R122Q$^{4.40}$, and G279S$^{7.44}$ were activated to a similar activation level as wild type hA$_1$AR with pEC$_{50}$ values of 9.24 ± 0.08, 9.04 ± 0.14, and 9.27 ± 0.09, which were also not significantly different from the pEC$_{50}$ value of the wild type receptor. Mutant receptor A52V$^{2.47}$ had a much lower efficacy (1.86 ± 0.14) in the presence of 1 µM of CPA than wild type hA$_1$AR, and also showed a more than 400-fold decreased potency. The activation level of mutant receptor W188C$^{5.46}$ was similar to wild type hA$_1$AR (4.35 ± 0.10), while the potency of CPA was decreased by 10-fold.
The two mutant receptors with increased constitutive activity, namely H78L\textsuperscript{3.23} and S246T\textsuperscript{6.47}, also showed increased constitutive activity in concentration–growth curves. In response to CPA, mutant receptor S246T\textsuperscript{6.47} was activated to a similar \(E_{\text{max}}\) level (4.81 \(\pm\) 0.26) with a similar pEC\(_{50}\) value for CPA (9.42 \(\pm\) 0.33) as at the wild type hA\(_1\)AR (Figure 2C and Table 2). Interestingly, mutant receptor H78L\textsuperscript{3.23} showed a 4.5-fold increase in constitutive activity compared to wild type, where further activation could not be obtained anymore by the addition of CPA (Figure 2B and Table 2).

Next, we investigated whether the increased constitutive activity of these two mutants could be decreased using an inverse agonist, DPCPX (Figure 3). For mutant receptor S246T\textsuperscript{6.47}, DPCPX reduced the constitutive activity to wild type hA\(_1\)AR levels with a pIC\(_{50}\) value of 8.55 \(\pm\) 0.25. However, the high constitutive activity of mutant receptor H78L\textsuperscript{3.23} was not reduced by DPCPX.

![Figure 3](image-url). Concentration–inhibition curves of the hA\(_1\)AR inverse agonist DPCPX at the wild type A\(_1\)AR and the CAMs, H78L\textsuperscript{3.23}, and S246T\textsuperscript{6.47}. Data were normalized as ratio over basal activity of wild type hA\(_1\)AR (dotted line). Combined graphs are shown as mean \(\pm\) SEM from at least three individual experiments performed in duplicate. Data for wild type are shown in dark blue and for CAMs in green.

Evaluating the diverse pharmacological effects of these mutant receptors, we characterized mutant receptors H78L\textsuperscript{3.23} and S246T\textsuperscript{6.47} as CAMs, mutant receptors A20T\textsuperscript{1.43}, A52V\textsuperscript{2.47}, R122Q\textsuperscript{4.40}, W188C\textsuperscript{5.46}, and G279S\textsuperscript{7.44} as CIMs, and mutant receptors D55V\textsuperscript{2.50}, D55G\textsuperscript{2.50}, P86L\textsuperscript{3.31}, L134F\textsuperscript{4.52}, T257P\textsuperscript{6.58}, and S267I\textsuperscript{7.32} as loss of function mutants (LFMs) (Table 2).

2.4. Ligand Binding on Wild Type and Mutant hA\(_1\)ARs

Selected mutants with diverse effects on receptor activation, i.e., H78L\textsuperscript{3.23}, L134F\textsuperscript{4.52}, W188C\textsuperscript{5.46}, S246T\textsuperscript{6.47}, and G279S\textsuperscript{7.44}, were further investigated on ligand binding in a mammalian expression system. Wild type and mutant receptors were transiently transfected into Chinese Hamster Ovary (CHO) cells, and receptor expression levels were measured by ELISA. All mutant receptors were expressed on the cell surface with similar levels to the wild type hA\(_1\)AR (Figure 4A).

Affinity values of the radioligand [\(^3\)H]DPCPX and \(B_{\text{max}}\) values of wild type and mutant hA\(_1\)ARs were determined by homologous competition displacement assays on transiently transfected membranes (Figure 4 and Table 3). [\(^3\)H]DPCPX had a pKD value of 8.36 \(\pm\) 0.03 at the wild type hA\(_1\)AR, which was significantly higher than the value on LFM L134F\textsuperscript{4.52} (8.06 \(\pm\) 0.08), but lower than the value on CIM G279S\textsuperscript{7.44} (8.62 \(\pm\) 0.06, Table 3). Mutant receptors H78L\textsuperscript{3.23}, W188C\textsuperscript{5.46}, and S246T\textsuperscript{6.47} showed similar pKD values of [\(^3\)H]DPCPX compared to the wild type hA\(_1\)AR. Diverse \(B_{\text{max}}\) values were obtained on mutant receptors in comparison to wild type hA\(_1\)AR (1.18 \(\pm\) 0.14 pmol/mg). A significantly increased expression level of 3.74 \(\pm\) 0.65 pmol/mg was observed on LFM L134F\textsuperscript{4.52}, while expression levels of CAMs H78L\textsuperscript{3.23} and S246T\textsuperscript{6.47} were decreased (0.17 \(\pm\) 0.01 pmol/mg).
and 0.11 ± 0.01 pmol/mg). Note that these values did not correlate with the cell surface expression data obtained from ELISA.

Figure 4. (A) Cell surface expression levels of wild type and mutant hA1AR transiently transfected on CHO cell membranes, as determined by ELISA. Data were normalized as ratio over mock-transfected CHO cells (mock, dotted line) and shown as mean ± SEM obtained from three individual experiments performed in pentaplicate. (B, C) Displacement of specific [3H]DPCPX binding to the transiently transfected wild type hA1AR, LFM L134F, CIMs W188C and G279S, and CAMs H78L and S246T on CHO cell membranes by DPCPX and CPA, respectively. Combined graphs are shown as mean ± SEM from three individual experiments, each performed in duplicate. Data for wild type are shown in dark blue, for CIMs shown in red, for CAMs in green, and for LFMs in grey.

Table 3. Affinity and B$_{\text{max}}$ values of [3H]DPCPX and binding affinity of CPA on wild type and mutant hA1ARs.

| Mutation  | [3H]DPCPX a pK$_D$ | [3H]DPCPX a B$_{\text{max}}$ (pmol/mg) | CPA pK$_i$ (High) | CPA pK$_i$ (Low) | Fraction (High) |
|-----------|-------------------|---------------------------------|----------------|----------------|-----------------|
| Wild type | 8.36 ± 0.03       | 1.81 ± 0.14                     | 9.24 ± 0.26    | 6.76 ± 0.05    | 0.15 ± 0.03     |
| H78L.3.23 | 8.46 ± 0.03       | 0.17 ± 0.01 **                 | 8.97 ± 0.35    | 6.83 ± 0.09    | 0.33 ± 0.04     |
| L134F.4.52| 8.06 ± 0.08 **    | 3.74 ± 0.65 **                 | 8.38 ± 0.29    | 6.26 ± 0.11 ** | 0.34 ± 0.03     |
| W188C    | 8.42 ± 0.03       | 1.87 ± 0.12                    | 8.02 ± 0.16 *  | 6.15 ± 0.01 ***| 0.29 ± 0.01     |
| S246T.6.47| 8.44 ± 0.05       | 0.11 ± 0.01 **                 | 8.98 ± 0.16    | 7.19 ± 0.08 ** | 0.26 ± 0.03     |
| G279S.4.44| 8.62 ± 0.06 *     | 2.11 ± 0.07                    | 8.74 ± 0.48    | 6.78 ± 0.06    | 0.17 ± 0.04     |

All values are shown as mean ± SEM obtained from at least three individual experiments performed in duplicate. a Values obtained from homologous displacement of ~1.6, 4.5, and 10 nM [3H]DPCPX from transiently transfected wild type and mutant CHO-hA1AR membranes at 25 °C. * p < 0.05; ** p < 0.01; *** p < 0.001 compared to wild type hA1AR, determined by one-way ANOVA with Dunnett’s post-test. Heterologous displacement by CPA of [3H]DPCPX radioligand binding on all mutant receptors as well as wild type hA1AR was best fitted to a two-site model (Figure 4C and Table 3). Wild type hA1AR had a pK$_i$ value of 9.24 ± 0.26 for the high-affinity state, 6.76 ± 0.05 for the low-affinity state with a fraction value of 0.15 ± 0.03 for the high-affinity state. Decreased pK$_i$ values were observed on CIM W188C for both high and low-affinity states (8.02 ± 0.16 at high affinity state and 6.15 ± 0.01 at low-affinity state). LFM L134F also showed a decreased affinity value of 6.26 ± 0.11 at the low-affinity state compared to the wild type receptor, while the high-affinity state was unchanged. Lastly, CAM S246T had an increased affinity value of 7.19 ± 0.08 at the low-affinity state with an unaffected affinity in the high-affinity state.
2.5. [\textsuperscript{35}S]GTP\textgamma{}S Functional Assay on Wild Type and Mutant hA\textsubscript{1}ARs

CHO cell membranes transiently transfected with wild type and mutant hA\textsubscript{1}AR were further tested in a functional assay, i.e., GTP\textgamma{}S binding (Figure 5 and Table 4). All selected mutant receptors showed similar basal activity to wild type hA\textsubscript{1}AR. In response to CPA, wild type hA\textsubscript{1}AR showed a potency/pEC\textsubscript{50} value of 8.98 ± 0.08 and an E\textsubscript{max} value (ratio over wild type basal activity) of 1.48 ± 0.13. Only CIM W188C\textsuperscript{5.46} showed altered receptor pharmacology upon activation by CPA with a decreased potency value of 8.28 ± 0.10, while the efficacy was not significantly affected. While LFM L134F\textsuperscript{4.52} did not show any activation in the yeast system, it could be activated in the mammalian system with similar potency and efficacy values for CPA compared to wild type. CAM S246T\textsuperscript{6.47} showed altered receptor pharmacology upon CPA-mediated activation with a higher pEC\textsubscript{50} value of 9.44 ± 0.22 and a slightly lower efficacy value of 1.21 ± 0.10 than wild type hA\textsubscript{1}AR, albeit not significantly different. CIM G279S\textsuperscript{7.44} did not show significantly different receptor pharmacology to wild type hA\textsubscript{1}AR in the mammalian system.

Table 4. Potency and efficacy values of CPA and DPCPX in [\textsuperscript{35}S]GTP\textgamma{}S binding assays on wild type and mutant hA\textsubscript{1}ARs.

| Mutation | Basal \textsuperscript{a} | CPA | DPCPX |
|----------|----------------|------|------|
|          | pEC\textsubscript{50} (−log M) | E\textsubscript{max} \textsuperscript{a} | pIC\textsubscript{50} (−log M) | I\textsubscript{max} \textsuperscript{b} |
| Wild type | 1.00 ± 0.09 | 8.98 ± 0.08 | 1.48 ± 0.13 | 8.09 ± 0.16 | 0.67 ± 0.05 |
| H78L\textsuperscript{3.23} | 1.24 ± 0.10 | 9.09 ± 0.12 | 1.40 ± 0.10 | 8.19 ± 0.25 | 0.83 ± 0.03 |
| L134F\textsuperscript{4.52} | 1.12 ± 0.17 | 9.08 ± 0.16 | 1.48 ± 0.24 | 8.14 ± 0.23 | 0.68 ± 0.01 |
| W188C\textsuperscript{5.46} | 1.21 ± 0.06 | 8.28 ± 0.10 \textsuperscript{*} | 1.94 ± 0.02 | 7.87 ± 0.25 | 0.74 ± 0.03 |
| S246T\textsuperscript{6.47} | 1.08 ± 0.10 | 9.44 ± 0.22 | 1.21 ± 0.10 | 8.44 ± 0.10 | 0.70 ± 0.05 |
| G279S\textsuperscript{7.44} | 1.17 ± 0.13 | 8.69 ± 0.10 | 1.57 ± 0.20 | 8.23 ± 0.06 | 0.65 ± 0.08 |

All values are shown as mean ± SEM obtained from at least three individual experiments performed in duplicate. \textsuperscript{a} Values were calculated as ratio over basal activity of wild type hA\textsubscript{1}AR. \textsuperscript{b} Values were calculated as ratio over basal activity of wild type or mutant hA\textsubscript{1}AR. \textsuperscript{*} p < 0.05 compared to wild type hA\textsubscript{1}AR, determined by one-way ANOVA with Dunnett’s post-test.

Next, we investigated whether the agonist-mediated activation could be inhibited by the antagonist, DPCPX, on wild type and mutant receptors (Figure 5B). For the wild type receptor, the activation level was reduced to 0.67 ± 0.05 with a pIC\textsubscript{50} value of 8.09 ± 0.16 for DPCPX. In the mammalian system, the CPA-mediated activation for all mutant receptors was reduced to wild type levels with similar pIC\textsubscript{50} values (Table 4).
2.6. Structural Mapping and Bioinformatics Analysis of Mutations

The mutations investigated in this study were mapped on the inactive (5UEN) and active (6D9H) hA1AR structure to provide structural hypotheses for the observed pharmacological effects (i.e., CIM, CAM, and LFM) of the different mutations, and explain differences between yeast and mammalian data. Mutations were found scattered over the receptor structure, with LFMs indicated in black, CIMs in red, and CAMs in green (Figure 6A). Whilst some LFMs can be considered drastic changes (for instance, T257P\textsuperscript{6.58} and P86L\textsuperscript{3.31}), others are relatively mild from a structural perspective (e.g., S267I\textsuperscript{7.32}). LFMs D55V/G\textsuperscript{2.50} sit in the sodium ion binding pocket in direct contact with the sodium ion (Figure 6B). The CAM S246T\textsuperscript{6.47} is found near the middle of helix 6, which undergoes a large conformational change upon the receptor activation (Figure 6C). Finally, W188C\textsuperscript{5.46} and L134F\textsuperscript{4.52} are positioned close to one another and point toward the membrane.

Figure 6. (A) Overview of all mutations mapped on the X-ray structure of the hA1AR, inactive (5UEN) in red and active (6D9H) in green. Residues are colored by their observed effect, CAMs in green, CIMs in red, and LFMs in black. (B) Close up of residue D55\textsuperscript{2.50}. In grey, residues are found in the A2A AR binding site, with the sodium ion from that structure (PDB: 4EIY) in purple. (C) Residue S246\textsuperscript{6.47} is found near the hinging region of TM6, the outward motion of which is associated with receptor activation (shown with arrow). (D) Residues L134\textsuperscript{4.52} and W188\textsuperscript{5.46} form a cluster and are pointing toward the membrane.
3. Discussion

Although the role of hA₁AR in cancer progression remains unclear, a growing amount of studies suggest that hA₁AR is involved in cancer development [13,14]. Previous structural studies and crystal structures of hA₁AR provided us with information on crucial residues for ligand binding and receptor activation, as well as essential interactions in the inactive receptor state and G protein coupling [17,23–25]. Moreover, compared to other residues, accumulation of cancer-related mutations has been observed in highly conserved residues of the TM domains [26]. Therefore, in this study, we studied 13 single-site point mutations located at the 7-TM domains of A₁AR obtained from The Cancer Genome Atlas (TCGA). All mutations were examined in the S. cerevisiae system and selected mutations were further investigated in the mammalian system to improve our understanding of the mechanism of receptor activation with respect to cancer development and progression.

3.1. Mutations Located at the Top Part of the Receptor

Mutant receptors H78L³.23, P86L³.31, T257P⁶.58, and S267I⁷.32, located at the top, extracellular part of the receptor, all showed dramatic changes upon receptor activation in the yeast system. Mutant receptor H78L³.23 showed an extremely high constitutive activity, which could not be further induced by CPA or reduced by DPCPX (Figures 2C and 3 and Table 2). Although this could not be confirmed in the mammalian system (probably due to its low expression level), it indicates that H78L³.23-hA₁AR is locked in an active conformation, which has been described previously on mutant receptor G14T¹.37 in hA₁AR [27].

Similar expression levels were not observed between ELISA and homologous competition assays (Figure 4A and Table 3) due to different experimental setups that whole-cell expressions of functioning receptors were determined in homologous competition assays [28]. Crystallographic structural evidence of the inactive state A₁AR reveals that H78³.23 forms a salt bridge with E164, which is important for the stabilization of a β-sheet between EL1 and EL2 [24]. It had been shown in a previous study on A₁AR that mutant receptor P86F³.31 resulted in abolished CPA binding. This indicates that the proline at residue 86 indirectly affects ligand binding by reorienting the TM1 conformation to favor N₆ substituents [30]. Both P86L³.31 and P86F³.31 are mutations in which the small size and rigid residue proline were exchanged by larger amino acids with hydrophobic side chains. The introduction of these larger side chains is potentially the causal factor for the loss of receptor activation. The residue T257⁶.58, located at the top part of the helix 6, forms a hydrophobic pocket along with M177⁵.35, L253⁶.54, and T270⁷.35, which has been shown to accommodate the antagonist DU172 in the A₁AR [24]. In A₂A AR, an alanine mutation at residue T256⁶.58 has been shown to result in decreased affinity of the reference antagonist ZM241385 [31].

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3.2. Mutations Located on Conserved Residues

Conserved residues and motifs of GPCRs are known to mediate ligand binding and receptor functionality [19]. Thus, mutations located at these residues may cause prominent alterations in receptor pharmacology. Alanine at residue 2.47 is highly conserved among class A GPCRs (72%) [33]. Mutant receptor A52V showed a dramatic decrease in both potency and efficacy of CPA (Figure 2A and Table 2), which could not be confirmed in mammalian cells due to a lack of expression. Interestingly, this same mutation occurs in CCR5, where this seemingly small change in the side chain, has been reported to greatly affect the binding of CCL5 [34], indicating the essential role of residue A2.47 in receptor–ligand interaction.

Two LFMs, D55G and D55V, are found at residue D2.50, which is the most highly conserved residue among class A GPCRs (92%) [35]. Mutations at residue D2.50 are known to alter ligand binding and/or G protein signaling [37,38]. Abolished G protein signaling has also been reported on mutant receptor D52N in A2A AR, in which it was shown that inter-helical packing was impacted by the change from aspartic acid to asparagine [37]. Therefore, our results implicate that the loss of the negatively-charged side chain in D2.50 impedes electrostatic interactions with Na+-ions and, thereby, leads to decreased receptor activation.

S246 belongs to the conserved CWxP motif in helix 6, which is classified as the microswitch region and associated with receptor activation [39]. In the CWxP motif, cysteine at residue 6.47 is conserved by 71% among class A GPCRs and serine is 10% [35]. In both yeast and mammalian systems, mutant receptor S246T showed slightly increased potency values of CPA (Figures 2C and 5A, Tables 2 and 4). The increase in potency value could be caused by the increase in ligand binding of CPA (Figure 4C). Of note, the affinity of DPCPX was unchanged, implying that this antagonist has no preference for binding to the active or inactive state of the receptor [40]. Additionally, hA1AR was not locked in the active conformation by mutation S246T, as DPCPX could still deactivate the receptor (Figure 3). Similarly, in the β2-adrenergic receptor, the mutation C285T has been characterized as a CAM, while C285S had similar properties to the wild type receptor [39]. As it is known that residue 6.47 is crucial for the rotamer toggle switch [39], a threonine mutation on 6.47 may alter the side chain modulation of the rotamer toggle switch, further impacting the movement of TM6 during receptor activation.

3.3. Mutations Located on Residues Pointing towards the Membrane

In mammalian cell membranes, cholesterol has been reported to have a modulatory role in the GPCR function via interaction with residues in the lipid–protein interface [41]. Moreover, compared to the membranes of mammalian cells, the yeast cell membrane contains less cholesterol and more ergosterol, which may result in a different receptor conformation and, thus, the functionality of human GPCRs between expression systems [41,42]. Moreover, the conflicting results obtained from different expression systems could be caused by differences in receptor expression levels.

Mutant receptor G279S has been characterized as a CIM with retained potency and efficacy of CPA in the yeast system, while decreased constitutive activity could not be observed in the mammalian system, possibly due to the slightly higher expression level than wild type hA1AR. Interestingly, G279S has also been identified as a Parkinson’s disease-associated mutation, which did not alter receptor expression or ligand binding but influenced the heteromerization with the dopamine D1 receptor [43].

Mutant receptor W188C showed a 10-fold decrease in the potency value of CPA in both yeast and mammalian systems (Figures 2C and 5A, Tables 2 and 4). This decrease in potency was caused by the decrease in the affinity of CPA (Figure 4C and Table 3). Despite the maintenance of hydrophobicity of the side chain, the substitution of tryptophan for cysteine introduced a dramatic reduction in the side chain size. Reducing the amino acid side chain size at position W188 may affect the receptor–ligand interaction of CPA on hA1AR. Moreover, it has been shown that W188 together with residues V137 and...
F144^{4.62}, W146, Y182^{5.40}, F183^{5.41}, and V187^{5.45} are part of a hydrophobic core, which, along with residues S150 and R154, forms contacts with the EL2 of two A_{1}AR homodimers in mammalian cells [24]. It has been hypothesized that EL2 exerts a crucial role in the transition between G protein-coupled and -uncoupled states [44]. While it was previously suggested that A_{1}AR homodimerizes, leading to cooperative orthosteric ligand binding in mammalian cells [45], the homodimerization of A_{1}AR in yeast cells remains undetermined.

Residue L134^{4.52} forms a cluster with W188^{5.46} pointing towards the membrane (Figure 6D). Mutant receptor L134F^{4.52} has been characterized as LFM in the yeast system. However, it behaved quite similar to wild type A_{1}AR in the mammalian system (Figure 5 and Table 4). L134^{4.52} is conserved amongst all ARs and is located close to the highly conserved residue in TM4, W^{4.50}. The latter is known to be involved in ligand binding and interaction with the cell membrane via cholesterol, where complete loss of ligand binding has been observed previously by mutating tryptophan to other amino acids [41,46,47]. Phenylalanine mutation at L134^{4.52} might thus indirectly change the interactions among residues W132^{4.50}, L99^{3.44}, A100^{3.45}, L193^{5.51}, and Y200^{5.58} [46], by the dramatic size change of the side chain, and this might be different when using a different cell membrane background.

3.4. Potential Role for hA_{1}AR Mutations in Cancer

Activation of hA_{1}AR has been identified with anti-proliferative effects in colon cancer, glioblastoma, and leukemia [10,48,49]. Mutations with inhibitory effects on receptor activation identified from colon cancer, such as the LFM D55G^{2.50} and CIM W188C^{5.46}, might then behave as pro-proliferative regulators in cancer progression. In contrast, deletion or blockade of hA_{1}AR resulted in inhibited cell proliferation but induced PD-L1 upregulation in melanoma cells, which led to compromised anti-tumor immunity [50]. Additionally, the hA_{1}AR antagonist DPCPX shows inhibitory effects on tumor cell proliferation and migration while promoting apoptosis [12,15]. Mutant receptors with the altered binding affinity of DPCPX, namely L134F^{4.52} and W188C^{5.46} in this study, may thus impact the efficacy of DPCPX treatments. Of note, due to the low frequency in comparison to known driver mutations in cancer patients, these cancer-related mutations in hA_{1}AR are unlikely to be cancer drivers [51]. However, passenger mutations should not be ruled out in consideration of cancer personalized therapy [52].

4. Materials and Methods

4.1. Data Mining

Mutation data were downloaded from The Cancer Genome Atlas (TCGA, version August 8, 2015; note, the TCGA data were frozen in early 2016) by using the Firehose tool [53]. MutSig 2.0 data were extracted when available, MutSig 2CV was used in cases where the former was not available. sequence data were filtered for missense somatic mutations and the A_{1}AR (Uniprot identifier P30542). The GPCRdb alignment tool was used to assign Ballesteros Weinstein numbers [22,55] to the positions through which a selection could be made for transmembrane domain positions.

4.2. Materials

The MMY24 strain and the S. cerevisiae expression vectors, the pDT-PGK plasmid and the pDT-PGK_hA_{1}AR plasmid (i.e., expressing the wild type receptor) were kindly provided by Dr. Simon Dowell from GSK (Stevenage, UK). The pcDNA3.1(+) plasmid cloned with N-terminal 3xHA-tagged hA_{1}AR was ordered from cDNA Resource Center (Bloomsburg, PA, USA). The QuikChange II® Site-Directed Mutagenesis Kit containing XL10-Gold ultracompetent cells was purchased from Agilent Technologies (Amstelveen, The Netherlands). The QIAprep mini plasmid purification kit and QIAGEN® plasmid
midi kit were purchased from QIAGEN (Amsterdam, The Netherlands). Adenosine deaminase (ADA), 1,4-dithiothreitol (DTT), 8-cyclopentyl-1,3-dipropylxanthine (DPCPX), and 3-amino-[1,2,4]-triazole (3-AT) were purchased from Sigma-Aldrich (Zwijndrecht, The Netherlands). N6-cyclpentyladenosine (CPA) was purchased from Santa Cruz Biotechnology (Heidelberg, Germany). Bicinchoninic acid (BCA) and BCA protein assay reagent were obtained from Pierce Chemical Company (Rockford, IL, USA). Radioligands 1,3-[3H]-dipropyl-8-cyclopentylxanthine ([3H]DPCPX, specific activity of 137 Ci × mmol−1) and [35S]-Guanosine 5′-(γ-thio)triphosphate ([35S]GTPγS, a specific activity 1250 Ci × mmol−1) were purchased from PerkinElmer, Inc. (Waltham, MA, USA). Rabbit anti-HA antibody (71-5500) was purchased from Thermo Fisher Scientific (Waltham, MA, USA), while goat anti-rabbit IgG HRP was purchased from Jackson ImmunoResearch Laboratories (West Grove, PA, USA).

4.3. Generation of hA1AR Mutations

The plasmids carrying hA1AR mutations were constructed by polymerase chain reaction (PCR) mutagenesis as previously described, using pDT-PGK_hA1AR or pcDNA3.1_hA1AR with N-terminal 3xHA tag as the template [17]. The QuikChange Primer Design Program of Agilent Technologies (Santa Clara, CA, USA) was used to design primers for mutant receptors, and primers were purchased from Eurogentec (Maastricht, The Netherlands). All DNA sequences were verified by Sanger sequencing at LGTC (Leiden, The Netherlands).

4.4. Transformation in MMY24 S. Cerevisiae Strain

The plasmids, pDT-PGK_hA1AR, containing either wild type or mutated hA1AR were transformed into a MMY24 S. cerevisiae strain following the lithium acetate procedure [56].

4.5. Liquid Growth Assay

In order to characterize the mutant hA1ARs, liquid growth assays in 96-well plates were performed to obtain concentration–growth curves as previously described [17]. Briefly, yeast cells expressing wild type or mutant hA1AR were inoculated to 1 mL selective YNB medium lacking uracil and leucine (YNB-UL) and incubated overnight at 30 °C. The overnight cultures were then diluted to 40,000 cells/mL (OD600 ≈ 0.02) in a selective medium without uracil, leucine, and histidine (YNB-ULH). For the determination of constitutive activity, 50 μL of yeast cells and 150 μL of YNB-ULH medium containing different concentrations of 3-AT and 0.8 IU/mL ADA were then added to each well. To obtain concentration–growth curves, 2 μL of various concentrations of ligands, 50 μL of yeast cells, and 150 μL of YNB-ULH medium containing 7 mM 3-AT and 0.8 IU/mL ADA were then added to each well. After incubation at 30 °C for 35 h in a Genios plate reader (TECAN, Zürich, Switzerland) with shaking for 1 min at 300 rpm every 10 min, the optical density was measured at a wavelength of 595 nm, which represented the level of yeast cell growth.

4.6. Cell Culture, Transient Transfection, and Membrane Preparation

Chinese hamster ovary (CHO) cells were cultured at 37 °C in 5% CO2 in a Dulbecco’s Modified Eagle’s Medium/Ham’s F12 (1:1, DMEM/F12) containing 10% bovine calf serum, streptomycin (50 μg/mL), and penicillin (50 IU/mL). Cells were grown until 80–90% confluency and subcultured twice weekly.

Transient transfection of CHO cells with wild type or mutated hA1AR plasmid constructs was performed using a polyethylenimine (PEI) method [57]. Cells were seeded in 10-cm culture dishes to achieve 50–60% confluency 24 h prior to transfection. On the day of transfection, cells were transfected with a PEI: DNA ratio of 3:1 and a plasmid DNA amount of 10 μg/dish. Moreover, 24 h post-transfection, the medium was refreshed, and 48 h after transfection, cells were collected and membranes were prepared as previously described [58]. Membranes were aliquoted in 250 or 100 μL and stored at −80 °C. Membrane protein concentrations were determined using the BCA method [59].
4.7. Enzyme-Linked Immunosorbent Assay

The ELISA experiments were performed with some modifications from a previously published procedure [60]. Moreover, 24 h after transfection, cells were seeded in a 96-well plate with a density of 10^6 cells per well; 48 h post-transfection, the cells were fixed with 4% formaldehyde and blocked with 2% bovine serum albumin (BSA) (Sigma-Aldrich Chemie N.V., Zwijndrecht, The Netherlands) in Tris-buffered saline (TBS) for 1 h. Then, the cells were incubated with rabbit anti-HA tag primary antibody (1:2500) in TBST (0.05% Tween 20 in TBS) overnight at 4 °C. The cells were washed 3 times in TBST and incubated with the goat anti-rabbit IgG HRP secondary antibody (1:6000) for 1 h at RT. After removing the secondary antibody and washing the cells with TBS, 3,3',5,5'-tetramethyl-benzidine (TMB) was added and incubated for 10 min in the dark. The reaction was stopped with 1 M H_3PO_4, and absorbance was read at 450 nm using a Wallac EnVision 2104 Multilabel reader (PerkinElmer).

4.8. Radioligand Displacement Assay

The displacement assays were performed as described previously [27]. Briefly, to each well the following was added: 25 µL cell membrane suspension, 25 µL of 1.6 nM radioligand [3H]DPCPX, 25 µL of assay buffer (50 mM Tris-HCl, pH 7.4), and 25 µL of six increasing concentrations of DPCPX (10^{-11} to 10^{-6} M) or CPA (10^{-10} to 10^{-5} M), all dissolved in assay buffer. Note, the number of cell membranes (10–25 µg) was adjusted to obtain (approximately) a 1500 DPM assay window for each mutant. Nonspecific binding was determined in the presence of 10^{-4} M CPA and represented less than 10% of the total binding. For homologous competition assays, radioligand displacement experiments were performed in the presence of 3 different concentrations of [3H]DPCPX (1.6, 4.5, and 10 nM) as well as 6 increasing concentrations of DPCPX (10^{-11} to 10^{-6} M). Incubations were terminated after 1 h at 25 °C by rapid vacuum filtration through GF/B filter plates (PerkinElmer, Groningen, Netherlands) using a Perkin Elmer Filtermate-harvester. Afterward, filter plates were washed ten times with ice-cold buffer (50 mM Tris-HCl, pH 7.4) and dried at 55 °C for 30 min. After the addition of 25 µL per well of the Microscint scintillation cocktail (PerkinElmer, Groningen, The Netherlands), the filter-bound radioactivity was measured by scintillation spectrometry in a Microbeta2® 2450 microplate counter (PerkinElmer).

4.9. [35S]GTPγS Binding Assay

[35S]GTPγS binding assays were adapted from a previously published method [27]. Membrane aliquots containing 15 µg protein were incubated with a total volume of 80 µL of assay buffer (50 mM Tris-HCl buffer, 5 mM MgCl_2, 1 mM EDTA, 100 mM NaCl, 0.05% BSA, and 1 mM DTT pH 7.4 supplemented with 10 µM GDP and 10 µg saponin) and 9 increasing concentrations of CPA (10^{-11} to 10^{-6} M) or 9 increasing concentrations of DPCPX (10^{-11} to 10^{-6} M) in the presence of a fixed concentration (EC_{80} for wild type or mutant hA_1ARs) of CPA for 30 min at 4 °C. Then, 20 µL of [35S]GTPγS (final concentration of 0.3 nM) was added to each well, followed by 90 min of incubation at 25 °C. Incubations were terminated and filter-bound radioactivity was measured as described above.

4.10. Modelling

Structures of the A_1AR in the inactive (PDB: 5UEN) [24] and active (PDB: 6D9H) states [23] and the inactive state of the A_2AAR (PDB: 4EIY) [38] were retrieved from the PDB. Missing side chains and loop regions were added using the GPCR-ModSim web server [61]. All structures were aligned to the inactive A_1AR, and figures were generated using the PyMOL Molecular Graphics System version 2.0 (Schrödinger, LLC., New York, NY, USA).

4.11. Data Analysis

All experimental data were analyzed by GraphPad Prism 7.0 or 8.0 (GraphPad Software Inc., San Diego, CA, USA). Data from yeast liquid growth and [35S]GTPγS binding
assays were analyzed by non-linear regression using “log (agonist) vs. response (three parameters)” or “log (inhibitor) vs. response (three parameters)” to obtain potency (EC\textsubscript{50}), inhibitory potency (IC\textsubscript{50}), and efficacy (E\textsubscript{\text{max}} or I\textsubscript{\text{max}}) values. The radioligand displacement curves were obtained from a statistically preferred one-site or two-site binding model. pK\textsubscript{i} values were calculated from pIC\textsubscript{50} values using the Cheng–Prusoff equation, where K\textsubscript{D} values were obtained from the homologous competition assays from this study and calculated by non-linear regression using “one site–homologous” [62].

5. Conclusions

In conclusion, 13 cancer-induced somatic mutations located at the 7-transmembrane domain of the adenosine A\textsubscript{1} receptor were retrieved from TCGA and characterized in a robust yeast system. Moreover, two CAMs (H78L\textsuperscript{3.23} and S246T\textsuperscript{6.47}), one LFM (L134F\textsuperscript{4.52}), and two CIMs (W188C\textsuperscript{5.46} and G279S\textsuperscript{7.44}) were also investigated in mammalian cells. The yeast system is a simplified, suitable, rapid, and accurate method for initial mutation screening that enables us to identify mutations with a dramatic effect on receptor activation. However, the current study shows that this system is best used for receptor mutations on the extracellular side, ligand-binding pocket, or pointing inwards from the membrane. Based on the results of this study, follow-up studies focusing on the effects of these mutations on other G protein coupling pathways, as well as in a disease-relevant system, are warranted to further investigate the effect of these hA\textsubscript{1}AR mutations in cell proliferation and migration, and eventually in cancer progression. Taken together, this study will enrich our understanding of the largely undefined role of hA\textsubscript{1}AR in cancer progression, which may eventually improve cancer therapies.

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