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PEGylated isoprenaline reveals distinct functions of cardiac β-adrenergic receptors located in the T-tubule vs. outer surface membrane

Marion Barthé¹, Flora Lefebvre², Emilie Langlois³, Florence Lefebvre¹, Patrick Lechène¹, Xavier Iturrioz³, Tâp Ha-Duong⁴, Laurence Moine², Nicolas Tsapis², Rodolphe Fischmeister¹,*

¹Université Paris-Saclay, Inserm, UMR-S 1180, Châtenay-Malabry, France.
²Université Paris-Saclay, CNRS, Institut Galien Paris-Saclay, Châtenay-Malabry, France.
³Center for Interdisciplinary Research in Biology (CIRB), Collège de France, Paris, France.
⁴Université Paris-Saclay, CNRS, BioCIS, Châtenay-Malabry, France.

*Corresponding author:
Rodolphe Fischmeister, PhD
INSERM UMR-S 1180
Faculty of Pharmacy, Université Paris-Saclay
5, rue JB Clément
92296 Châtenay-Malabry
France
Tel: +33 1 46 83 57 71
Fax: +33 1 46 83 54 75
E-mail: rodolphe.fischmeister@inserm.fr
Abstract

Membrane proteins are present in both cardiac T-tubule (TTM) and outer surface membrane (OSM), although at a different density. Classical pharmacology does not allow to explore the function of a membrane protein separately in OSM vs. TTM. Here, we developed a technology based on size exclusion to explore the function of β-adrenergic receptors (β-ARs) located in the OSM. We synthetized a PEG-Iso molecule by covalent linking between isoprenaline (Iso) and a 5000 Da PolyEthylene-Glycol (PEG). Using confocal microscopy, we show that PEGylation constrains molecules outside the T-tubule network. PEG-Iso produced similar effects as Iso on $I_{Ca_{L}}$, sarcomere shortening and $Ca^{2+}$ transients. However, PEG-Iso increased [cAMP], with a lower efficacy than Iso, produced a much lower stimulation of nuclear PKA activity than Iso but a larger stimulation of cytosolic PKA at equivalent levels of [cAMP]. Our results show that activation of OSM β-ARs is sufficient to activate cytosolic PKA and excitation-contraction coupling, but insufficient to activate nuclear PKA or nuclear protein phosphorylation for which additional activation of TTM β-ARs is needed.
Introduction

The β-adrenergic receptor (β-AR) is a key player in the regulation of cardiac function during sympathetic nerve stimulation. The classical pathway for β-AR receptor signalling is activation of adenylyl cyclases (AC) via Gαs, resulting in increased intracellular cAMP levels ([cAMP]i)\(^1\). The primary target of cAMP is the cAMP-dependent protein kinase (PKA) that in turn phosphorylates several key proteins involved in the excitation-contraction (EC) coupling, such as the L-type Ca\(^{2+}\) channel (LTCC or Ca\(_{v1.2}\)), phospholamban, troponin I, etc.\(^2\).

The phosphorylation of Ca\(_{v1.2}\) (or its regulatory protein Rad\(^3,4\)) leads to enhanced LTCC current (I\(_{Ca,L}\)) and sarcoplasmic reticulum (SR) Ca\(^{2+}\) release via the ryanodine type 2 receptor (RyR2), contributing to enhanced Ca\(^{2+}\) transients and contraction\(^5,6\). Phosphorylation of phospholamban increases Ca\(^{2+}\) uptake into the SR which accelerates Ca\(^{2+}\) transient decay and, together with troponin I phosphorylation, speeds up relaxation. Where as short term stimulation of β-AR/cAMP is beneficial for the heart, chronic activation of this pathway results in altered Ca\(^{2+}\) signalling, cardiac hypertrophy and fibrosis, leading to ventricular dysfunction\(^7\) and cardiac arrhythmias\(^8-12\).

The cell membrane of cardiomyocytes is characterized by invaginations of the surface membrane, occurring primarily perpendicular to myocyte longitudinal edges, at intervals of ~1.8–2 µm, that form a complex interconnected tubular network penetrating deep into the cell interior\(^13\). This network is called the transverse (T-) tubules (TT) system, although the invaginations may often bifurcate in the axial direction or form branches\(^13\). These tubular structures are found mostly in adult ventricular myocytes, where they represent about 30% of the total cell membrane\(^14,15\), and occur near the sarcomeric z-discs\(^16\), allowing functional junction with the SR called dyad. They are critical for EC coupling by concentrating LTCCs and
positioning them at close proximity of RyR2 clusters at the junction of SR to form Ca$^{2+}$ release units$^{17,18}$. During an action potential, TT propagate the cell-membrane depolarization inside the cell allowing Ca$^{2+}$ entry to trigger successive Ca$^{2+}$ release thus promoting synchronicity as well as efficiency of the Ca$^{2+}$-induced Ca$^{2+}$ release (CICR) phenomenon$^{19}$. Their enrichment with ion channels and the presence of signaling pathways components such as β-ARs$^{20}$ and ACs$^{21}$ make these structures essential for cardiomyocyte function and regulation.

Biochemical assays following membrane fractionation have provided indirect evidence that membrane proteins may have different properties whether located in TTM or OSM. Other indirect evidence came from experiments in cardiomyocytes in which TTM was uncoupled from OSM (detubulation) using a hyperosmotic shock with molar concentrations of formamide$^{14,22-30}$. However, none of these approaches allowed to investigate in an intact cardiomyocyte whether the function of a given membrane receptor differs if the receptor is located in OSM or TTM. To do so requires to being able to separately activate or inhibit the receptor in OSM or TTM which has not been feasible so far. Here, we tackled this challenge by developing a size exclusion strategy using the PEGylation technology. By a covalent link between isoprenaline (Iso) and a PolyEthylene-Glycol (PEG) chain, we increased the size of the β-AR agonist to prevent it from accessing the TT network. Our working hypothesis is that PEGylated isoprenaline (PEG-Iso) activates only β-ARs present in OSM while free Iso activates β-ARs present in both OSM and TTM. We thus characterized the properties of PEG-Iso and compared its functional effects in adult rat ventricular myocytes (ARVMs) with those of Iso.
Results

**PEGylation constrains molecules outside the TT network.** The rationale for using ligand PEGylation as a tool to impede ligand diffusion into T-tubules came from the observation that fluorescent PEG5000 molecules were unable to access to TTs. A typical experiment (out of 15) using confocal microscopy is shown in Fig. 1. When ARVMs were exposed during 15 min to 100 µM of PEG5000 molecules functionalized with Fluorescein Isothiocyanate (FITC), no fluorescence was seen throughout the cell (Fig. 1a, b). On the contrary, when the cells were exposed during 15 min to 100 µM fluorescein, fluorescence was seen with clear staining of the TTs (Fig. 1c, d). We suspected that the lack of access of PEG-FITC in the TTs was due to steric hindrance due to the presence of glycocalyx matrix. To test this hypothesis, ARVMs were first exposed during 1 h to a solution containing 0.25 U/mL neuraminidase, an enzyme that degrades sialic acid in the glycocalyx matrix. As shown in Fig. 1e, f, this allowed PEG-FITC to enter the TTs as evidenced by the striated profile now observed.

**PEG-Isoprenaline binds to β₁- and β₂-adrenergic receptors.** The above experiments demonstrate that PEG5000 molecules are unable to access the TTs in our experimental conditions. We thus synthesized PEGylated isoprenaline (PEG-Iso) by linking isoprenaline to PEG5000 molecules (Supplementary Fig. 1). Our rationale was that since PEG-Iso does not access TTs, it will not reach β-ARs located in the TTM. But the question remained whether PEG-Iso would be able to bind to β-ARs located in the OSM. To address this question, we performed radioligand binding studies in purified membranes from Chinese Hamster Ovary (CHO) cells overexpressing either β₁- or β₂-ARs. Competition curves between \[^{125}\text{Iodo}\text{cyanopindolol}\text{ and }\text{Iso}\text{ or PEG-Iso were used to measure }K_i\text{ values for both ligands.}\]
As shown in Fig. 2a, b, PEG-Iso binds to both β₁- and β₂-ARs, but with an affinity which is ~2-orders of magnitude lower than Iso. We hypothesized that the decreased affinity of PEG-Iso could be due to wrapping of the PEG chain around Iso moiety reducing its exposure to solvent (water). To test this, we used molecular dynamics simulations to evaluate the solvent-accessible surface area (SASA) of Iso moiety in PEG-Iso compared to free Iso. Fig. 2c shows that free Iso SASA remains steady over the whole studied time range (black line). In contrast, Iso moiety in PEG-Iso (red line) has a SASA which largely fluctuates and reaches many times values close to zero, indicating its frequent shielding by the PEG chain. We calculated that only 1.2% of PEG-Iso conformations have an Iso SASA larger than 90% of the free Iso average value. This means that over a time lapse of 100 s, the Iso moiety is unwrapped from the PEG chain and therefore able to bind to a receptor during only a short time of about 1 s.

**Comparison of the effects of PEG-Iso and Iso on cytosolic cAMP.** The next series of experiments were designed to test whether PEG-Iso was able to produce a functional β-AR response in ARVMs. First, cytosolic cAMP ([cAMP]ᵢ) was monitored in isolated ARVMs expressing the FRET-based sensor Epac-S²¹⁸⁷. As seen in Fig. 3, both Iso (Fig. 3a, b) and PEG-Iso (Fig. 3c, d) produced a concentration-dependent increase in [cAMP]ᵢ. However, there were two major differences: 1) the concentration-response curve to PEG-Iso was shifted ~100-fold towards larger concentrations as compared to Iso; 2) the maximal efficacy of PEG-Iso was significantly lower by ~30% than that of Iso (Fig. 3e). The former was anticipated based on the lower binding affinity of PEG-Iso to β-ARs shown above. The latter can be explained by the fact that PEG-Iso only activates β-ARs in OSM while Iso activates β-ARs in both OSM and TTM, thus producing a larger [cAMP]ᵢ elevation.
Comparison of the effects of PEG-Iso and Iso on $I_{\text{Ca,L}}$. Next, the ability of Iso and PEG-Iso to stimulate the L-type $Ca^{2+}$ current, $I_{\text{Ca,L}}$, was compared at a single concentration of each, i.e. 10 nM Iso and 1 µM PEG-Iso, shown above to produce an equivalent response on $[\text{cAMP}]_i$ (Fig. 4). As shown in the individual experiments in Fig. 4a, b, both Iso and PEG-Iso produced an increase in the amplitude of $I_{\text{Ca,L}}$ and both responses were on average similar in amplitude (Fig. 4c).

Comparison of the effects of PEG-Iso and Iso on excitation-contraction coupling. To investigate the impact of a stimulation of OSM β-ARs on EC coupling, $Ca^{2+}$ transients and sarcomere shortening were simultaneously recorded in Fura-2-loaded ARVMs and their response to PEG-Iso and Iso were evaluated. Here, two concentrations of PEG-Iso were used, 100 and 300 nM, and two concentrations of Iso producing similar elevations of $[\text{cAMP}]_i$, 1 and 3 nM, respectively. As shown in Fig. 5 and 6, 100 and 300 nM PEG-Iso increased contractility (Fig. 5b), $Ca^{2+}$ transients (Fig. 6b) and accelerated their relaxation kinetics (Fig. 5d and 6d). These effects were not different from those produced by 1 and 3 nM Iso, respectively, on either sarcomere shortening (Fig. 5a, c), $Ca^{2+}$ transient amplitude (Fig. 6a, c) or relaxation kinetics (Fig. 5c, d; Fig. 6c, d).

Comparison of the effects of PEG-Iso and Iso on cytosolic and nuclear PKA activity. The observation that equipotent concentrations of PEG-Iso and Iso on $[\text{cAMP}]_i$ produced equivalent effects on $I_{\text{Ca,L}}$ and EC coupling, may suggest that OSM β-ARs are not functionally different from their TTM homologs as long as their degree of activation leads to similar $[\text{cAMP}]_i$ responses. However, the above experiments do not exclude possible differences in
the compartmentalization of intracellular cAMP cascade when cAMP is synthesized upon OSM or TTM β-AR stimulation. To obtain further insight, we compared the effect of PEG-Iso and Iso on PKA activity in the bulk cytoplasm and in the nucleus. For that, we used genetically encoded A-kinase activity FRET-based reporters (AKAR3) targeted to these compartments by the addition of a nuclear export sequence (NES), and a nuclear localizing sequence (NLS), respectively. As shown previously, adenoviral transfer allowed robust and compartment-specific expression of these biosensors after 24 h in ARVMs. In the experiments shown in Fig. 7, four increasing concentrations of Iso (0.3, 1, 3 and 10 nM: Fig. 7a, c) and PEG-Iso (10, 30, 100 nM and 1 µM: Fig. 7b, d) were successively applied to ARVMs infected with AKAR3-NES probe and cytosolic PKA activation was measured. Although higher concentrations of ligands could not be used because of the saturation of the AKAR3 probe, both compounds produced a clear concentration-dependent increase in cytosolic PKA activity, with a similar maximal effect but a ~100-fold rightward shift in concentrations for the effect of PEG-Iso as compared to Iso. However, when nuclear PKA activity was measured, a clear difference between the effects of Iso and PEG-Iso were observed (Fig. 8): While PEG-Iso and Iso still produced a concentration-dependent increase in nuclear PKA activity, the maximal response to PEG-Iso (Fig. 8d, e) was reduced about 2-fold as compared to Iso (Fig. 8b, e). The same difference is observed regarding the nuclear proteins phosphorylation levels (Supplementary Fig. 2) after stimulation with Iso 10 nM or PEG-Iso 1 µM.

Supplementary Fig. 3 shows a plot of cytosolic and nuclear PKA activity as a function of cytosolic cAMP from the average data in the experiments in Fig. 3, 7 and 8 for 3 concentrations of PEG-Iso (10 and 100 nM and 1 µM) and Iso (0.3, 1 and 10 nM). It shows that for any given measured increase in [cAMP]i, PEG-Iso is more efficient than Iso to
increase PKA activity in the cytosol (Supplementary Fig. 3a), while on the contrary Iso is more efficient than PEG-Iso to increase PKA activity in the nucleus (Supplementary Fig. 3b).

**Discussion**

The main function of the TT system is to provide proximity between LTCCs in the TTM and RyR2 in the SR membrane\(^\text{18,35}\). However, TTM also contains many other membrane proteins, such as receptors, enzymes, ion channels and transporters. These include plasma membrane \(\text{Ca}^{2+}-\text{ATPase}\)\(^\text{36}\), \(\text{Na}^{+}-\text{Ca}^{2+}\) exchanger\(^\text{37,38}\), \(\text{Na}^{+}\) channels\(^\text{37}\), \(\text{Na}^{+}/\text{K}^{+}-\text{ATPase}\)\(^\text{39}\), ACs\(^\text{21,40}\) and \(\beta\)-ARs\(^\text{20,41}\). Density of these membrane proteins is usually found to be higher in TTM than in the external membrane. Hence, the TT network plays a determinant role in rapid activation and synchronous \(\text{Ca}^{2+}\) release, cellular signalling and, consequently, cardiac contraction in both human and animal models\(^\text{42-44}\).

Any given membrane protein is likely to serve different functions and be regulated in a different manner whether it is located in TTM or in OSM. Attempts to address this important question have so far been based on the elimination of the TT network (detubulation) using a hyperosmotic shock with molar concentrations of formamide\(^\text{14,22-30}\). This technique, first used in skeletal muscle\(^\text{45}\), was introduced in the cardiac field by Kawai et al.\(^\text{30}\) and provided an excellent experimental tool for numerous T-tubular studies. However, the potential role of shock-induced detubulation in physiologically and pathophysiological relevant conditions is essentially unknown. The method is very harsh on the cells as less than 10% of the cells survive to the procedure, which raises issues about whether the "survivors" are representative cells\(^\text{15}\). Besides, osmotic shock produces a sealing of the TT which remain
present as vesicles inside the cell\textsuperscript{22} and may still respond to hormonal or pharmacological challenges even if they are electrically disconnected from the surface membrane\textsuperscript{46}.

To overcome these limitations, we introduce here a completely different approach based on size exclusion. We proposed to enlarge the size of a ligand molecule by attaching it to a chain of PolyEthylene Glycol (PEG) so that it does not access TTM but remains active on OSM. PEG is a hydrophilic, flexible and rather inert polymer. The covalent linking of one or several PEG chains to a therapeutic molecule, called PEGylation\textsuperscript{47}, is commonly used in several products on the market\textsuperscript{48}. PEGylated molecules exhibit improved biodistribution and pharmacokinetics, better stability and solubility, reduced immunogenicity and longer plasma half-life due to both reduced renal filtration and proteolysis, when compared to non-PEGylated analogues\textsuperscript{49,50}. In our study, we diverted the same technology towards an entirely different objective: instead of using PEGylated molecules to improve drug formulation, pharmacokinetics and efficacy, we propose to use PEGylated drugs as key holders to prevent drug (key) access in the TT network and thus limit its access to the outer surface of the cell. We provide a proof of concept that this approach works using PEG\textsubscript{5000} functionalized with non-permeant FITC (PEG-FITC): When ARVMs were exposed to PEG-FITC, fluorescence was only seen on the periphery of the cell while when the non-PEGylated free diffusible dye was used, fluorescence was seen in the TT compartment (Fig. 1). Thus, large molecular weight PEGs are prevented to diffuse within TTS whereas small molecular weight fluorophore can easily enter.

This finding was surprising, though, since longitudinal TT diameter from healthy control ARVMs varies from 50 to 350 nm (\textasciitilde200 nm on average\textsuperscript{51,43,51}), which is quite larger than the size of a PEG\textsubscript{5000} molecule on the order of 5 nm. However, a number of recent studies have shown that solute movement in the TT network is strongly restricted even
though the diameter of TTs is far larger than the molecular dimensions of typical solutes.\textsuperscript{52-56}

Impediment of solute movement in the TT network was proposed to be caused by the presence of the glycocalyx\textsuperscript{57} at the cell surface, reducing the effective diameter of a TT\textsuperscript{53, 54, 58}. This was supported by our finding that pre-treatment of isolated ARVMs with neuraminidase, an enzyme which cleaves sialic acid from oligosaccharide chains in the glycoprotein matrix on the external cell surface, allowed PEG-FITC to enter the TT compartment (Fig. 1). Thus, when the protocol used for cell isolation is sufficiently gentle to preserve the integrity of the extracellular matrix in the TT, which must be the case in our experimental conditions, it is possible to prevent a ligand from accessing the TTM if the size of the ligand molecule is sufficiently enlarged. We selected PEG molecules between 3000 and 5000 Da MW which represents a good compromise between solubility issues and size exclusion.

PEG-Iso was thus synthetized by linking isoprenaline to PEG\textsubscript{5000} molecules. Characterization of its binding properties showed that PEG-Iso binds to β\textsubscript{1}- and β\textsubscript{2}-ARs but with \textasciitilde{}2-orders of magnitude lower affinity than Iso (Fig. 2). Since PEG was grafted on the lateral amino group of Iso (Supplementary Fig. 1), a position shown to maintain its affinity for β-ARs\textsuperscript{59}, it was unlikely that the chemical modification \textit{per se} was the reason for the decreased affinity. We thus explored a possible involvement of the PEG conformation in this phenomenon using molecular dynamics simulations. Because of its hydrophilicity and flexibility, PEG is very dynamic in solution and constantly changes conformation. Consequently, the Iso moiety in PEG-Iso is rarely fully accessible to solvent and spends most of the time with the PEG chain wrapped around it (Fig. 2). Based on the calculations, only \textasciitilde{}1\% of PEG-Iso conformations are able to bind β-ARs as compared to 100\% for free Iso,
which is in the order of magnitude of the measured difference in their respective binding affinity on β-ARs.

When applied on ARVMs, PEG-Iso produced stimulatory effects on [cAMP]ᵢ, PKA activity, Iₒᵤ, sarcomere shortening and Ca²⁺ transients which all had the hallmarks of a β-AR stimulation (Fig. 3 to 8). Because of the reduced binding affinity towards β-ARs, PEG-Iso produced its effect at ~2-orders of magnitude larger concentrations than Iso. But there were two other striking differences between the effects of PEG-Iso and Iso which strongly suggest that the two ligands did not act on the exact same populations of receptors.

First, PEG-Iso produced a stimulation of [cAMP]ᵢ with ~30% lower maximal efficacy than Iso (Fig. 3). The simplest explanation for this result, which is what drove us to undertake this study, is that PEG-Iso activates only β-ARs located in the OSM while Iso activates β-ARs located in both OSM and TTM. Since both populations of β-ARs are likely coupled to ACs, one would expect an activation of OSM β-ARs with PEG-Iso to lead to a smaller maximal increase in [cAMP]ᵢ than an activation of all β-ARs with Iso. This diminution is in line with decreased cAMP production sites by detubulation. Cyclic AMP must emanate mainly from sarcolemmal β₁-ARs, which produce highly diffusible signals throughout the cell in contrast to β₂-ARs. However, there are conflicting results regarding the distribution of β-ARs on the cardiac cell membrane. Immunohistochemical data have suggested that β₁- and β₂-ARs are present in OSM and TTM in mouse heart. Using radioligand binding, β₁-AR density was found to be almost 2-fold more concentrated in OSM than TTM, whereas β₂-AR density was evenly distributed across the entire cell surface in dog heart. Using nanoscale live-cell scanning ion conductance and fluorescence resonance energy transfer microscopy techniques in healthy ARVMs, Nikolaev et al. found that β₁-ARs are distributed across the entire cell surface while β₂-ARs are localized exclusively in TTM. While the distribution of
β₁-ARs across the entire cell surface was confirmed in ARVMs in another study using formamide detubulation, β₂-ARs were found to be only present in OSM\textsuperscript{23}. Additional experiments using PEGylated selective β₁- and β₂-AR agonists and antagonists are needed to solve this issue and to precisely characterize the role of each β-AR subtype in the different membrane compartments.

Second, PEG-Iso produced a much lower stimulation of nuclear PKA activity than Iso (Fig. 8) even though both ligands were equally efficient in stimulating cytosolic PKA activity (Fig. 7). Also, when cytosolic and nuclear PKA activity measured during PEG-Iso or Iso stimulation were plotted as a function of [cAMP] measured under the same conditions, PEG-Iso was found to be more efficient than Iso to increase PKA activity in the cytosol, while on the contrary Iso is more efficient than PEG-Iso to increase PKA activity in the nucleus (Supplementary Fig. 3). A likely interpretation of these results is that β-ARs located in TTM (or a β-AR subtype more concentrated in TTM) are more efficiently coupled to nuclear PKA than those located in OSM, while β-ARs located in OSM (or a β-AR subtype more concentrated in OSM) are more efficiently coupled to cytosolic PKA. In a recent study, we showed that β₂-ARs, unlike β₁-ARs, are inefficient in activating nuclear PKA activity in ARVMs\textsuperscript{33}. Thus, if the ratio of β₂/β₁ receptors is larger in OSM than in TTM, as discussed above\textsuperscript{23}, then this would explain the lower efficacy of PEG-Iso to activate nuclear PKA activity as compared to Iso. Recent studies have shown that perinuclear PKA has the ability to activate without detachment of the catalytic subunits, and thus to phosphorylate its targets in close proximity\textsuperscript{63, 64}. According to these studies, only supra-physiological concentrations of cAMP could lead to dissociation of the catalytic subunits and their translocation into the nucleus. Perinuclear PKA is anchored to the nuclear membrane via its interaction with mAKAP and its dynamics are finely regulated by ACS and PDE4D\textsuperscript{35}, ACS being concentrated
in TTM\textsuperscript{21} and PDE4D3 being linked to mAKAP\textsuperscript{66}. The location of AC5 may therefore be crucial for nuclear signalling and explain why β-ARs present in TTM are necessary for activation of nuclear PKA.

Another important finding from this study concerns the contribution of OSM β-ARs to the regulation of EC coupling. Activation of β-ARs only in OSM with PEG-Iso produced a similar increase in $I_{Ca,L}$ amplitude, sarcomere shortening and Ca\textsuperscript{2+} transients as Iso, when both ligands were used at concentrations producing an equivalent elevation of [cAMP].

Since OSM contributes to ~60% of total cell membrane in ARVMs\textsuperscript{15}, either β-ARs and ACs are more concentrated in OSM than TTM, or they are in large excess over what is needed to activate PKA phosphorylation of proteins involved in EC coupling. Also, cAMP produced at OSM must diffuse rapidly in the cytosol in order to activate PKA phosphorylation of substrates located deep inside the cell, such as LTCCs in TTM. A recent study showed that under basal conditions, cAMP is bound to proteins and its diffusion in the cytosol is slow; but when cAMP increases in the cytosol, e.g. as a result of β-AR stimulation, the cAMP binding sites become saturated and cAMP diffuses freely\textsuperscript{67}.

While this study provides new insight on the differential function of OSM vs. TTM β-ARs in healthy cardiomyocytes, it raises obvious questions on how their role is altered in cardiac pathology. A reduction in the density and a disorganization of the TT network was observed in pathological conditions\textsuperscript{17, 51}, with great impact on EC coupling and contractile function\textsuperscript{18}. A previous study showed a redistribution of β\textsubscript{2}-ARs from TTM to OSM in failing ARVMs\textsuperscript{20}. As β-ARs are not simply bystanders but also participate in the remodeling during pathological hypertrophy and failure, it is important to know how the OSM vs. TTM subpopulations of β-AR subtypes contribute to this process. The PEGylation technology should allow to elucidate the changes that operate in OSM vs. TTM distribution of β-ARs in
ventricular cells isolated from diseased rats, and to determine how this impacts on cellular function.

In conclusion, this study provides a novel approach to distinguish the function of a membrane protein depending on its location on the cell membrane. Whereas we focused here on the function of β-ARs, the size exclusion strategy provided by ligand PEGylation can be extended to other ligands, such as selective agonists or antagonists of β-AR subtypes or other G-protein coupled receptors, dihydropyridine agonists or antagonists of LTCCs, etc. More generally, it should pave the road to exploring and comparing the function of any membrane receptor, channel, transporter or enzyme in TTM and OSM compartments in intact cardiomyocytes.

Methods

Animals. All animal care and experimental procedures complied with the ARRIVE guidelines and conform to the European Community guiding principles in the Care and Use of Animals (Directive 2010/63/EU of the European Parliament), the local Ethics Committee (CREEA Ile-de-France Sud) guidelines, and the French decree no. 2013-118 of February 1st, 2013 on the protection of animals used for scientific purposes (JORF no. 0032, February 7th, 2013, p2199, text no. 24). Animal experiments according were carried out according to the European Community guiding principles in the care and use of animals (2010/63/UE), the local Ethics Committee (CREEA Ile-de-France Sud) guidelines and the French decree n° 2013-118 on the protection of animals used for scientific purposes.
Synthesis of PEG-ISO. Isoprenaline was grafted into the reactive end of 5000 Da PEG (PEG_{5000}). A carbodiimide reaction was carried out between the carboxylic acid of PEG and the amine function of Iso (Supplementary Fig. 1). In a 25 mL amber vial, 41.3 mg of DCC (0.2 mmol; 2eq), 23 mg of NHS (0.2 mmol; 2eq) and 500 mg of OH-PEG_{5000}-COOH (0.1 mmol; 1eq) were solubilized in 10 mL of dichloromethane. 1 mL of triethylamine (TEA) was added to the syringe. The solution was stirred for 5 h at RT. 21.1 mg isoprenaline (0.1 mmol; 1 eq) were added to the reaction mixture. The mixture was left to stir overnight. The solution was then filtered through silica gel and precipitated twice in cold diethyl ether. After evaporation of the ether, the final powder was solubilised in H₂O.

Purification of PEG-ISO. A purification step was introduced to obtain a pure PEG-Iso product and to eliminate the various by-products obtained, potential impurities and especially traces of ungrafted Iso which could interfere in the biological experiments. An HPLC 1290 Infinity II - (Agilent Technologies) consisting of a 4-channel binary pump, a UV/visible detector (PDA diode array), a 1260 Infinity DEDL light scattering detector and a C18 XBridge column, 4.6 x 150 mm, 5 µm (17 mL/min flow rate) was used. The mobile phase consisted of a mixture of water + 0.1% formic acid and acetonitrile. This phase was pushed according to a gradient from 1 to 100% in 15 min into the stationary phase for the purification of PEG-Iso. Detection was carried out in the UV at 254 and 280 nm. PEG-Iso was solubilized in water. Several successive injections of 300 µL each were performed. The retention time of PEG-Iso was 11 min. The fractions containing PEG-Iso were recovered, evaporated and then frozen and freeze-dried. Finally, the powder obtained was stored at -20°C and protected from light.
Cardiomyocyte isolation and culture. Adult male Wistar rats (250–300 g) were anesthetized by intraperitoneal injection of pentobarbital (0.1 mg/g) and hearts were excised rapidly. Individual adult rat ventricular cardiomyocytes were obtained by retrograde perfusion of the heart as previously described. For enzymatic dissociation, the hearts were perfused during 5 min at a constant flow of 6 mL/min at 37°C with a Ca²⁺-free Ringer solution containing (in mM): NaCl 117, KCl 5.7, NaHCO₃ 4.4, KH₂PO₄ 1.5, MgCl₂ 1.7, D-glucose 11.7, Na₂-phosphocreatine 10, taurine 20, and 4-(2-hydroxyethyl)piperazine-1-ethanesulfonic acid (HEPES) 21, pH 7.1. This was followed by a perfusion at 4 mL/min for 40 min with the same solution containing 1 mg/mL of collagenase A (Roche Diagnostics GmbH, Mannheim, Germany) plus 300 µM ethylene glycol tetraacetic acid (EGTA) and CaCl₂ to adjust free Ca²⁺ concentration to 20 µM. The ventricles were then separated, finely chopped and gently agitated to dissociate individual cells. The resulting cell suspension was filtered on gauze and the cells were allowed to settle down. The supernatant was discarded and cells resuspended four more times with Ringer solution at increasing [Ca²⁺] from 20 to 300 µM. Freshly isolated cells were suspended in minimal essential medium (MEM: M 4780; Sigma, St Louis, MO USA) containing 1.2 mM [Ca²⁺] supplemented with 2.5% foetal bovine serum (FBS, Invitrogen, Cergy-Pontoise, France), 1% penicillin–streptomycin (P-S), 20 mM HEPES (pH 7.6), and plated on 35 mm, laminin-coated (10 mg/mL) culture dishes at a density of 10⁴ cells per dish and kept at 37°C (5% CO₂). After 1 h, the medium was replaced by 300 µL of FBS-free MEM. To perform FRET imaging, the medium was replaced by 300 µL of FCS-free MEM or transduced with an adenovirus encoding the Epac-S₉₉₇ FRET-based sensor, or the AKAR3 FRET-based sensor addressed to the cytosol or the nucleus, at a multiplicity of infection (MOI) of 1000 pfu/cell and cells were cultivated for 36 h prior to the experiments. Patch-clamp and IonOptix experiments were performed on cells 24 h after dissociation.
**Confocal imaging.** Ventricular cardiomyocytes from adult rats were deposited on 35 mm glass bottom Petri dishes. Two hours after deposition, the control cells were brought into contact with commercial fluorescent PEG_{5000} functionalized with FITC (Fluorescein Isothiocyanate) (Nanocs) or free Fluorescein (Sigma-Aldrich) chosen as positive control. Other cells were treated with neuraminidase during 1 h at 0.25 U/mL. After that, cells were incubated with PEG-FITC. All molecules were dissolved in Ringer containing (in mM): NaCl 121.6, KCl 5.4, MgCl$_2$ 1.8, CaCl$_2$ 1.8, NaHCO$_3$ 4, NaH$_2$PO$_4$ 0.8, D-glucose 5, sodium pyruvate 5, HEPES 10, adjusted to pH 7.4. The acquisitions were made with a Leica TCS SP5 confocal microscope, a white light laser and an X40 oil immersion objective. The cells were excited at 495 nm and fluorescence was recovered at wavelengths >510 nm.

**Molecular dynamics simulations.** Molecular dynamics simulations of free Iso and PEG-Iso were carried out to predict their conformations in aqueous solution and in particular to assess the solvent-accessibility of Iso moiety once grafted to the PEG. A 3000 Da PEG molecule was used in these simulations. The 2D chemical structure of each of the two molecules was first converted into a three-dimensional structure using ChemAxon's MarvinSketch chemical editing software. The topology and GAFF force field parameters$^{69}$ used in this study were then automatically generated using the ACPYPE program$^{70}$. Initial three-dimensional structure was then placed in the centre of 13.4 nm side cubic simulation. These dimensions allow the solute not to interact with its virtually created images due to the periodic boundary conditions used to simulate an infinitely duplicated mesh. The box was then filled with water (model TIP3P$^{71}$) and 150 mM NaCl. The system was then subjected to 2 short simulations (1 ns each) to equilibrate first the temperature around 300 K and then
the pressure around 1 bar. The Nosé-Hoover\textsuperscript{72,73} (coupling time $\tau_T = 0.5$ ps) and Parrinello-Rahman\textsuperscript{74} ($\tau_P = 2.5$ ps) algorithms were used for maintaining constant temperature and pressure, respectively. Finally, each of the two molecules was subjected to a production run of 150 ns. All simulations were performed using the GROMACS software version 2016\textsuperscript{75}. To analyse the trajectories, we mainly used the GROMACS internal $gmx sasa$ tool to calculate the solvent-accessible surface area (SASA) for iso moiety in both free Iso or PEG-Iso molecules. To compare the solvent-accessibility of iso moiety in free Iso and PEG-Iso, the percentage of PEG-Iso conformations with a $SASA_{\text{PEG-Iso}} \geq 0.9$ $SASA_{\text{free-Iso}}$ was calculated.

**Binding experiments.** Binding assays were carried out in a final volume of 100 $\mu$L, containing membrane suspension from CHO cells overexpressing human $\beta_1$- or $\beta_2$-adrenergic receptors, 145 pM of the radioligand $[^{125}\text{I}]c$yanopindolol and non-radioactive ligands at concentrations ranging from $10^{-10}$ to $10^{-3}$ M. Incubation was carried out for 2 h at room temperature and terminated by addition of 4 mL PBS. Then, rapid filtration was performed through Whatman GF/C glass fiber filters previously soaked in PBS containing 0.33% polyethyleneimine using a compact cell harvester (Millipore\textsuperscript{®} 1225 Sampling Vacuum Manifold). Binding reaction was transferred to the filters, and washed 3 times with Wash Buffer. Filter-bound radioactivity was measured in a gamma counter. All determinations were performed at least in triplicates.

**FRET imaging.** FRET experiments were performed at room temperature 36 h after cell plating. Cells were maintained in a Ringer solution containing (in mM): NaCl 121.6, KCl 5.4, MgCl$_2$ 1.8; CaCl$_2$ 1.8; NaHCO$_3$ 4, NaH$_2$PO$_4$ 0.8, D-glucose 5, sodium pyruvate 5, HEPES 10, adjusted to pH 7.4. Images were captured every 4 s using the 40x oil immersion objective of
a Nikon TE 300 inverted microscope connected to a software-controlled (Metaflor, Molecular Devices, Sunnyvale, CA, USA) cooled charge coupled (CCD) camera (Sensicam PE, PCO, Kelheim, Germany). Cells were excited during 150-300 ms by a Xenon lamp (100 W, Nikon, Champigny-sur-Marne, France) using a 440/20BP filter and a 455LP dichroic mirror. Dual emission imaging was performed using an Optosplit II emission splitter (Cairn Research, Faversham, UK) equipped with a 495LP dichroic mirror and BP filters 470/30 (CFP) and 535/30 (YFP), respectively. Spectral bleed-through into the YFP channel was subtracted using the formula: $YFP_{corr} = YFP - 0.6 \times CFP$.

**Electrophysiological experiments.** The whole cell configuration of the patch-clamp technique was used to record $I_{Ca,L}$. Patch electrodes had resistance of 0.5–1.5 MΩ when filled with internal solution containing (in mM): CsCl 118, EGTA 5, MgCl$_2$ 4, Na$_2$-phosphocreatine 5, Na$_2$ATP 3.1, Na$_2$GTP 0.42, CaCl$_2$ 0.062, Hepes 10, adjusted to pH 7.3 with CsOH. External Cs$^+$-Ringer solution contained (in mM): NaCl 107.1, CsCl 20, NaHCO$_3$ 4, NaH$_2$PO$_4$ 0.8, D-glucose 5, sodium pyruvate 5, Hepes 10, MgCl$_2$ 1.8, CaCl$_2$ 1.8, adjusted to pH 7.4 with NaOH. The cells were depolarized every 8 s from −50 mV to 0 mV during 400 ms. The junction potential was adjusted to give zero current between pipette and bath solution before the cells were attached to obtain a tight gigascale (>1GΩ). The use of −50mV as holding potential allowed the inactivation of voltage-dependent sodium currents. Currents were not compensated for capacitance and leak currents. The amplitude of $I_{Ca,L}$ was measured as the difference between the peak inward current and the current at the end of the depolarization pulses. The cells were voltage-clamped a patch-clamp amplifier (either RK-400 (Bio-Logic, Claix, France) or Axopatch 200B (Axon Instruments, Inc., UnionCity, CA, USA)). Currents were analog filtered at 5 kHz and digitally sampled at 10 kHz using a 16-bit analog to digital
converter (DT321; Data translation, Marlboro, MA, USA or a Digidata1440A, Axon Instruments) connected to a PC (Dell, Austin, TX, USA).

**Measurements of Ca$^{2+}$ transients and sarcomere shortening.** All experiments were performed at room temperature. Isolated ARVMs were loaded with 1 μM Fura-2 AM (Thermo Fischer Scientific) and Pluronic acid (0.012%, Thermo Fischer Scientific) for 15 min in a Ringer solution containing (in mM): KCl 5.4, NaCl 121.6, sodium pyruvate 5, NaHCO$_3$ 4.013, NaH$_2$PO$_4$ 0.8, CaCl$_2$ 1.8, MgCl$_2$ 1.8, glucose 5, Heps 10, pH 7.4 with NaOH. Sarcomere shortening and Fura-2 ratio (measured at 512 nm upon excitation at 340 and 380 nm) were simultaneously recorded in Ringer solution, using a double excitation spectro fluorimeter coupled with a video detection system (Ion Optix, Milton, MA, USA). Myocytes were electrically stimulated, with biphasic field pulses (5 V, 4 ms) at a frequency of 1 Hz. Ca$^{2+}$ transient amplitude was measured by dividing the twitch amplitude (difference between the end-diastolic and the peak systolic ratios) by the end-diastolic ratio, thus corresponding to the percentage of variation in the Fura-2 ratio. Similarly, sarcomere shortening was assessed by its percentage variation, which was obtained by dividing the twitch amplitude (difference between the end-diastolic and the peak systolic sarcomere length) by the end-diastolic sarcomere length. Relaxation kinetics were estimated by a non-linear fit of the decaying part of the Ca$^{2+}$ transient and sarcomere shortening traces with the following equation: $X(t) = A \cdot \exp(-{(t-t_0)/\tau}) + A_0$, where $t_0$ is zero time, $A_0$ the asymptote of the exponential, $A$ the relative amplitude of the exponential and $\tau$ the time constant of relaxation. The maximum first derivative of transients during the deflexion allowed determination of the rising velocities of the signals. All parameters were calculated offline with dedicated software (IonWizard 6x, IonOptix R).
**Proteins expression analysis.** Cardiomyocytes were treated with or without isoprenaline 10 nM or PEG-Iso 1 µM (for 15 min) and the nuclei were enriched using a purification kit (proteoExtract). Nuclear proteins were examined by immunoblot for phosphorylation of PKA substrates (with a specific phospho-PKA antibody) and total protein. The ratio of phospho-PKA expression was assessed by densimetric scanning of immunoblots.

**Statistical analysis.** All results are expressed as mean±SEM. Statistical analysis was performed using GraphPad software (GraphPad Software, Inc., La Jolla, CA, USA). Normal distribution was tested by the Shapiro-Wilk normality test. For simple two-group comparison, we used an unpaired Student t test. Differences between multiple groups were analysed using an ordinary 1-way ANOVA with Tukey post hoc test. A P value<0.05 was considered statistically significant.

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Author contributions


Competing interests

The authors declare no competing interests.

Additional information

Supplementary information is available for this paper at:
Figure Legends

Fig. 1 Localization of PEG-FITC in freshly isolated adult rat ventricular cardiomyocytes.
Typical confocal images (a, c, e) and plot profiles representing the fluorescence intensity measured across the cell (b, d, f) of ARVMs incubated during 15 min with either 100 µM PEG5000-FITC (a, b), 100 µM free fluorescein (c, d), or 100 µM PEG5000-FITC after a 1h treatment with 0.25 U/mL neuraminidase (e, f).

Fig. 2 Binding properties of Iso and PEG-Iso on β-ARs. β1- or β2-AR CHO membrane preparations (2.5 µg/triplicate) were incubated for 2h with 0.25 nM [125I]iodocyanopindolol and increasing concentrations of unlabeled Iso or PEG-Iso. a Example of a typical competition binding of Iso and PEG-Iso on β1-ARs. b Example of a typical competition binding of Iso and PEG-Iso on β2-ARs. K_i values were, respectively, 2.0 ± 0.7 µM and 1.4 ± 0.7 mM for Iso and PEG-Iso on β1-ARs; 2.1 ± 0.4 µM and 429 ± 1 µM for Iso and PEG-Iso on β2-ARs (N=5). c Time evolution of solvent-accessible surface area (SASA) of Iso (black trace) and PEG-Iso (red trace) using molecular dynamics simulations. Two representative conformations of PEG-Iso with large or small SASA are shown.

Fig. 3 Effect of Iso and PEG-Iso on cytosolic cAMP in adult rat ventricular cardiomyocytes.
Freshly isolated ARVMs were infected with an adenovirus encoding the Epac-SH187 FRET-based cytosolic cAMP sensor for 48h at 37°C at a multiplicity of infection of 1000 pfu/cell. a Typical experiment showing the time course of CFP/YFP ratio during successive applications of four increasing Iso concentrations: 0.3, 1, 10 and 100 nM. b Similar experiment showing
the time course of CFP/YFP ratio during successive applications of four increasing PEG-Iso concentrations: 10 and 100 nM, 1 and 10 µM. Pseudo-color images shown above the main graphs were taken at times indicated by the corresponding letters in the graphs. c, d Summary data from several similar experiments as in a and b, respectively. The bars show the mean ± s.e.m of the data shown by symbols. 16 cells from 3 rats were used in c; 32 cells from 4 rats in d. One-way ANOVA and Tukey’s multiple comparisons post hoc test were used: ** p<0.01; *** p<0.001; ns, non significant. e Comparison of the concentration-response curves in c and d. One-way ANOVA and Tukey’s multiple comparisons post hoc test: *** p<0.001.

Fig. 4 Comparison of the effects of PEG-Iso and Iso on $I_{\text{Ca,L}}$. The whole-cell patch-clamp technique was applied to ARVMs after 24 h culture. a Typical experiment showing the time course of $I_{\text{Ca,L}}$ current density upon application of 10 nM Iso and during washout. b Similar experiment showing the response of $I_{\text{Ca,L}}$ current density to 1 µM PEG-Iso. Individual current traces shown above the graphs were taken at times indicated by the corresponding letters on the graphs. c Summary data from several similar experiments as in a and b. Student t-test: ns, non significant.

Fig. 5 Comparison of the effects of PEG-Iso and Iso on sarcomere shortening.

Representative traces of sarcomere shortening recorded in ARVMs paced at 0.5 Hz and loaded with Fura-2 AM (1 µM) showing the effect of Iso (1 and 3 nM, a) and PEG-Iso (100 and 300 nM, b). The bars in a and b show the mean ± s.e.m of the data shown by symbols. c, d Average time-to-50% relaxation of sarcomere shortening from experiments shown in a and
respectively. 30 cells from 4 rats were used in a and c; 34 cells from 4 rats in b and d. One-way ANOVA and Tukey’s post hoc test: * p<0.05; *** p<0.001; ns, non-significant.

Fig. 6 Comparison of the effects of PEG-Iso and Iso on Ca^{2+} transients. Representative traces of Ca^{2+} transients recorded in ARVMs paced at 0.5 Hz and loaded with Fura-2 AM (1 µM) showing the effect of Iso (1 and 3 nM, a) and PEG-Iso (100 and 300 nM, b). The bars in a and b show the mean ± s.e.m of the data shown by symbols. c, d Exponential time constant (Tau) of relaxation of Ca^{2+} transients from experiments shown in a and b, respectively. 30 cells from 4 rats were used in a and c; 34 cells from 4 rats in b and d. One-way ANOVA and Tukey’s post hoc test: * p<0.05; ** p<0.01; *** p<0.001; ns, non-significant.

Fig. 7 Comparison of the effects of PEG-Iso and Iso on cytosolic PKA activity. Freshly isolated ARVMs were infected with an adenovirus encoding the AKAR3-NES FRET-based PKA sensor for 48h at 37°C at a multiplicity of infection of 1000 pfu/cell. a, Typical experiment showing the time course of YFP/CFP ratio during successive applications of three increasing Iso concentrations: 0.3, 1 and 3 nM. b, Similar experiment showing the time course of YFP/CFP ratio during successive applications of three increasing PEG-Iso concentrations: 10, 30 and 100 nM. Pseudo-color images shown above the main graphs were taken at times indicated by the corresponding letters in the graphs. c, d Summary data from several similar experiments as in a and b, respectively. The bars show the mean ± s.e.m of the data shown by symbols. 3 rats and 8-10 cells were used in c; 3-5 rats and 13-18 cells in d. One-way ANOVA and Tukey’s multiple comparisons post hoc test: ** p<0.01; *** p<0.001. e Comparison of the concentration-response curves in c and d.
**Fig. 8 Comparison of the effects of PEG-Iso and Iso on nuclear PKA activity.** Freshly isolated ARVMs were infected with an adenovirus encoding the AKAR3-NLS FRET-based PKA sensor for 48h at 37°C at a multiplicity of infection of 1000 pfu/cell. **a,** Typical experiment showing the time course of YFP/CFP ratio during successive applications of three increasing Iso concentrations: 0.3, 1 and 3 nM. **b,** Similar experiment showing the time course of YFP/CFP ratio during successive applications of three increasing PEG-Iso concentrations: 10, 30 and 100 nM. Pseudo-color images shown above the main graphs were taken at times indicated by the corresponding letters in the graphs. **c, d** Summary data from several similar experiments as in **a** and **b,** respectively. The bars show the mean ± s.e.m of the data shown by symbols. 3 rats and 10-12 cells were used in **c**; 3-5 rats and 10-17 cells in **d.** One-way ANOVA and Tukey’s multiple comparisons post hoc test: * p<0.05; ** p<0.01; *** p<0.001. **e** Comparison of the concentration-response curves in **c** and **d.***
Figure 2
Figure 3
Figure 4

Calcium current density (pA/pF)

Iso 10 nM
PEG-isoo 1 μM

% increase in \( I_{Ca,L} \) amplitude
Figure 5
Figure 6
Figure 7
Figure 8