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Mitochondrial uncouplers induce proton leak by activating AAC and UCP1

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Abstract

Mitochondria generate heat due to H⁺ leak (I_H) across their inner membrane¹. I_H results from the action of long-chain fatty acids on uncoupling protein 1 (UCP1) in brown fat²⁻⁶ and ADP/ATP carrier (AAC) in other tissues^{1,7-9}, but the underlying mechanism is poorly understood. As evidence of pharmacological activators of I_H through UCP1 and AAC is lacking, I_H is

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Author contributions A.M.B., A.M.N., P.B., M.G. and Y.K. conceived the project and designed experiments. A.M.B. performed all of the electrophysiological experiments, except for pilot experiments and data for Fig. 1, which were performed by A.F.; A.M.B., E.T.C., L.K., J.H., T.B. and N.B. performed respirometry on C2C12 mitochondria. J.S. performed respiration experiments in C2C12 cells. A.M.N., P.B. and M.G. designed and performed computational analyses. A.M.B., A.M.N., P.B., M.G. and Y.K. wrote the manuscript. All of the authors discussed the results and commented on the manuscript.

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Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this paper.

Competing interests Y.K., M.G., P.B. and J.S. are shareholders of Equator Therapeutics. Y.K. is an advisor to Equator Therapeutics. Y.K. is shareholder of YourChoice Therapeutics. E.T.C. is a founder, board member and equity holder in Matchpoint Therapeutics. The other authors declare no competing interests.

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induced by protonophores such as 2,4-dinitrophenol (DNP) and cyanide-4-(trifluoromethoxy) phenylhydrazone (FCCP)^{10,11}. Although protonophores show potential in combating obesity, diabetes and fatty liver in animal models¹²⁻¹⁴, their clinical potential for treating human disease is limited due to indiscriminately increasing H⁺ conductance across all biological membranes^{10,11} and adverse side effects¹⁵. Here we report the direct measurement of I_H induced by DNP, FCCP and other common protonophores and find that it is dependent on AAC and UCP1. Using molecular structures of AAC, we perform a computational analysis to determine the binding sites for protonophores and long-chain fatty acids, and find that they overlap with the putative ADP/ATP-binding site. We also develop a mathematical model that proposes a mechanism of uncoupler-dependent I_H through AAC. Thus, common protonophoric uncouplers are synthetic activators of I_H through AAC and UCP1, paving the way for the development of new and more specific activators of these two central mediators of mitochondrial bioenergetics.

Mitochondrial I_H controls cellular bioenergetics by uncoupling H⁺ flow between the electron-transport chain and ATP synthase. I_H causes thermogenesis, increases body energy expenditure and limits mitochondrial production of reactive oxygen species^{1,16}. Pharmacological control of I_H is central to the development of effective treatments for metabolic and age-related disorders. Unfortunately, the choice of small-molecule activators of I_H through AAC and UCP1 is extremely limited. Physiologically, I_H through UCP1 and AAC is activated by free long-chain fatty acids (FAs)^{3,16-18}, but they cannot be used as drugs owing to their poor selectivity and bioavailability. Thus, protonophores such as DNP and FCCP have emerged as pharmacological tools to increase H⁺ conductance of the inner mitochondrial membrane (IMM). Protonophores are membrane-soluble weak acids that carry H⁺ across biological membranes without the need for membrane-transport proteins^{10,11} (Fig. 1e). This protein-independent mechanism of action is broadly accepted as universal for all biological and artificial lipid membranes, including the IMM.

Although DNP and FCCP have become indispensable tools for controlling energy metabolism at the mitochondrial, cellular and systemic levels, here we demonstrate that their mechanism of action was substantially misunderstood. Although these mitochondrial uncouplers do possess protonophoric activity, their major effect in the mitochondria is the activation of I_H through AAC and UCP1.

Preference of DNP and FCCP towards the IMM

We used the patch-clamp technique to directly record I_H induced by DNP and FCCP across the whole IMM of individual isolated mitochondria stripped of the outer membrane (mitoplasts)^{2,7,19,20}. Application of DNP and FCCP to the cytosolic side of skeletal muscle mitoplasts resulted in a high I_H . By contrast, H⁺ currents induced across the plasma membrane of HEK293 cells were considerably smaller even when much higher concentrations of DNP and FCCP were used (Fig. 1a-d). These first direct measurements of DNP- and FCCP-induced I_H across the IMM suggest that the IMM has a unique ability to potentiate the action of protonophores, either due to its distinct biophysical properties or the presence of specific membrane proteins.

Protonophores induce I_H through AAC

As AAC is responsible for FA-induced mitochondrial I_H in tissues other than brown/beige fat⁷, we hypothesized that DNP and FCCP might activate I_H through AAC. DNP induced I_H in a dose-dependent manner (Fig. 2a and Extended Data Fig. 1a). The AAC inhibitor carboxyatractyloside (CATR) suppressed the current by around 60% (Fig. 2a,c), whereas another AAC inhibitor, bongkreikic acid (BKA; which inhibits the m-state of AAC versus the c-state for CATR), had only a small effect (around 15% inhibition; Extended Data Fig. 1b).

There are two somatic AAC isoforms in mice, AAC1 and AAC2^{21,22}, with AAC1 being dominant in heart and skeletal muscle^{21,23}. Accordingly, I_H induced by DNP in cardiac IMMs from *AAC1*^{-/-} mice^{7,24} was reduced by around 80% (Fig. 2d,e) and was insensitive to CATR (Fig. 2b,c). Thus, DNP activates I_H across the IMM primarily by targeting AAC. The AAC-independent component of DNP-induced I_H (around 20%; Fig. 2d,e) may be associated with the protonophoric activity of DNP, its action on other IMM proteins or both.

We next investigated whether FCCP and other mitochondrial uncouplers also induce I_H through AAC. In agreement with previous studies^{11,25}, FCCP, SF6847^{11,25} and BAM15^{25,26} were more potent than DNP and induced robust mitochondrial I_H at nanomolar concentrations (Fig. 2g and Extended Data Fig. 2a,e). CATR inhibited I_H induced by FCCP, SF6847 and BAM15 by approximately 20% (Extended Data Fig. 1d, e), 50% (Extended Data Fig. 2a, c) and 70% (Extended Data Fig. 2e,g), respectively. I_H induced by FCCP, SF6847 and BAM15 in cardiac IMMs from *AAC1*^{-/-} mice was reduced by around 50% (Fig. 2f,g), 60% (Extended Data Fig. 2a,b,d) and 85% (Extended Data Fig. 2e,f,h), respectively. In contrast to the uncoupler-induced I_H , K^+ current activated across the IMM by the K^+ -selective ionophore valinomycin²⁷ was not reduced in cardiac IMMs from *AAC1*^{-/-} mice (Extended Data Fig. 2i-k). Interestingly, for all uncouplers, AAC1 deficiency had a larger effect on I_H than pharmacological inhibition with CATR (Fig. 2 and Extended Data Figs. 1 and 2). This could be due to competition between CATR and uncouplers for the AAC-binding site, or a CATR-insensitive AAC state induced by uncouplers (CATR binds only to the c-state of AAC²⁸).

We next compared the rates of respiration induced by uncouplers in mitochondria isolated from wild-type (WT) and *AAC1/AAC2* double-knockout (DKO) C2C12 cells⁷. DNP and BAM15 caused a dose-dependent increase in the rate of uncoupled respiration in mitochondria from WT C2C12 cells, which was substantially reduced in DKO mitochondria (Extended Data Fig. 3a-f).

We also compared uncoupled respiration induced by FCCP, DNP and BAM15 in intact WT and DKO C2C12 cells. DKO cells have a significantly lower basal respiration rate than WT cells⁷ (Extended Data Fig. 4a). ATP synthase inhibitor oligomycin reduced respiration of WT cells but not DKO cells, as they lack mitochondrial ATP production⁷ (Extended Data Fig. 4a). Further application of increasing FCCP concentrations stimulated uncoupled respiration in both WT and DKO cells, but DKO respiration was lower and plateaued at around 20 pmol s⁻¹ per 10⁶ cells (Extended Data Fig. 4b), indicating that the DKO cells have a limited respiratory capacity. The limited respiration capacity of

DKO cells was further confirmed using a combination of valinomycin and nigericin to induce AAC-independent mitochondrial uncoupled respiration (Extended Data Fig. 4f). As mitochondrial biomass and the expression of the electron-transport chain complexes was overall similar in DKO and WT cells (Extended Data Fig. 3g-m), reduced Krebs cycle activity or mitochondrial substrate availability in DKO cells (which fully rely on glycolysis for ATP production⁷) may limit maximal respiration.

Thus, to provide a valid comparison of uncoupler-induced respiration in WT and DKO cells, we analysed only lower concentrations of FCCP (100 nM to 1 μ M), DNP (10 μ M to 250 μ M) and BAM15 (500 nM to 5 μ M) that did not exceed the respiration ability of DKO cells (approximately 20 pmol s⁻¹ per 10⁶ cells). Respiration induced by all uncouplers was reduced in DKO cells, but FCCP showed an overall lower dependence on AAC compared with DNP and BAM15 (Extended Data Fig. 4b-e), correlating well with the electrophysiological data (Fig. 2 and Extended Data Fig. 2e-h). Moreover, as FCCP concentration increased, AAC dependence of the FCCP-induced respiration became even smaller (Extended Data Fig. 4e), probably due to the increased protonophoric component. By contrast, AAC dependence of BAM15-induced respiration remained constant (Extended Data Fig. 4e), perhaps due to its limited protonophoric activity²⁶.

Thus, in addition to their protonophoric activity, widely used mitochondrial uncouplers such as FCCP, DNP and BAM15 activate I_H through AAC. Notably, BAM15 is less toxic in cells and mice compared with FCCP and DNP^{29,30}, which may correlate with its lower protonophoric activity²⁶ and higher dependence on AAC (Extended Data Fig. 4e).

DNP and FCCP induce I_H through UCP1

Being the dominant uncoupling protein in brown and beige fat^{2-5,19,31}, UCP1 may be responsible for DNP- and FCCP-induced I_H in these tissues. IMM-associated phospholipases result in substantial amounts of endogenous FAs in the IMM of brown fat. Thus, DNP and FCCP were applied together with 10 mM methyl- β -cyclodextrin (M β CD) to extract endogenous FA and therefore limit their effect on UCP1^{2,19,20}. Higher concentrations of DNP and FCCP were applied while using M β CD because it binds to hydrophobic compounds non-specifically, reducing their effective concentrations. For example, 500 μ M DNP in the presence of 10 mM M β CD induces a similar I_H amplitude as 50 μ M DNP without M β CD in heart IMMs (Extended Data Fig. 1c).

Application of 500 μ M DNP together with 10 mM M β CD induced robust I_H in brown fat IMMs (Fig. 3a (left)). This current was strongly suppressed by 1 mM GDP (guanosine 5'-diphosphate), an established UCP1 inhibitor³¹ (Fig. 3a, b), and reduced in *UCP1*^{-/-} IMMs (Fig. 3a,c). Thus, UCP1 is crucial for DNP-induced I_H in brown fat mitochondria. Interestingly, DNP-induced I_H carried by UCP1 had a linear current-voltage curve (Fig. 3a (left)) compared with the inwardly rectifying current-voltage relationship of DNP-induced I_H carried by AAC in heart IMM, which showed a sharp increase in the amplitude of inward currents with membrane hyperpolarization (Fig. 2a and Extended Data Fig. 1c). In *UCP1*^{-/-} brown fat IMMs, the residual I_H was inwardly rectifying and GDP-insensitive (Fig. 3a,b), suggesting the involvement of AAC.

FCCP-induced I_H in brown fat IMM was also inhibited by GDP and significantly reduced in $UCP1^{-/-}$ mice (Fig. 3d,f). I_H remaining in $UCP1^{-/-}$ IMM was insensitive to GDP (Fig. 3d,e). Interestingly, FCCP-induced I_H was less dependent on UCP1 than DNP-induced I_H (~50% versus ~80% reduction in $UCP1^{-/-}$ IMM, respectively; Fig. 3c,f) similar to the relative AAC dependence of FCCP- and DNP-induced I_H in heart IMM (Fig. 2e,f). FCCP depends less on AAC and UCP1 than DNP due to either its higher protonophoric activity or the involvement of other IMM proteins.

Nucleotides inhibit DNP-induced I_H

Mg^{2+} -free ADP and ATP are negative physiological regulators of FA-induced I_H through UCP1 and AAC^{2,7,32-34}. Thus, we hypothesized that uncoupler-activated I_H is also inhibited by adenine nucleotides.

ATP inhibited I_H through UCP1 induced by 500 μ M DNP with a half-maximum inhibitory concentration (IC_{50}) of around 14 μ M, but the IC_{50} increased to about 80 μ M when 5 mM DNP was used (Extended Data Fig. 5a). Thus, ATP appears to compete with DNP for binding to UCP1, as previously observed for FA^{2,19}. As the nucleotide-binding site of UCP1 is not selective for ATP³¹, other purine nucleotides, such as GDP (Fig. 3a), also inhibit DNP-induced I_H through UCP1.

In contrast to UCP1, which binds only purine nucleotides on the cytosolic face of the IMM, AAC transports adenine nucleotides and binds them on both IMM faces, depending on the conformational state (c or m)²⁸. As the vast majority of AAC proteins in isolated mitoplasts are initially in the c-state, with the nucleotide-binding site exposed to the cytosolic face of the IMM⁷, 1 mM ADP was first applied only to the cytosolic (bath) solution. We observed a small, transient inhibition of DNP-induced I_H (Extended Data Fig. 5b), probably due to the binding of cytosolic ADP to the c-state of AAC, followed by a transition to the m-state and release of ADP to the matrix side. In this state, the translocation pathway is expected to become unobstructed as there is no ADP in the pipette solution, enabling DNP to rebind and allowing I_H to recover (Extended Data Fig. 5b).

We next applied ADP on both IMM sides to study how continuous nucleotide exchange interferes with I_H . The electroneutral ADP/ADP homoexchange was used to avoid I_H contamination with ADP/ATP heteroexchange current⁷. The addition of 1 mM ADP to both sides of the IMM led to a greater and persistent inhibition of DNP-induced I_H (compare Extended Data Fig. 5c with Extended Data Fig. 5b). This suggests that DNP cannot induce I_H in a fraction of AAC transporters that are nucleotide bound during continuous nucleotide exchange, either because of nucleotide–DNP competition or simple obstruction of the translocation pathway. As 1 mM ADP is at the higher end of physiologically relevant Mg^{2+} -free ADP concentrations, we propose that adenine nucleotides cannot completely suppress DNP-induced H^+ transport through AAC under physiological conditions but can downregulate it. The inhibitory effects of one-sided and two-sided ADP application on DNP-induced I_H are similar to those on FA-induced I_H through AAC⁷ and suggest that DNP and FA induce I_H after binding to the central translocation pathway of AAC and UCP1^{2,7}.

Uncouplers and FAs bind within AAC

We sought to identify putative binding sites for chemical uncouplers in AAC using in silico docking and molecular dynamics simulations. AAC1 is the only protein mediating I_H for which a high-resolution structure has been determined. In the c-state, AAC1 forms a large water-accessible cavity facing the cytosolic side of the IMM, which is occupied by CATR in the crystal structures³⁵⁻³⁷. Previous studies have identified the base of the cavity as the binding site for transported nucleotides^{38,39}. We targeted the entire cavity of bovine AAC1 (Protein Data Bank (PDB): 2C3E)³⁶ for molecular docking with the CATR molecule removed.

We identified highly reproducible binding modes for DNP deep in this cavity, about halfway across the IMM, which were unaffected by the protonation state of the compound (Fig. 4a-c). At this depth in the cavity, one side is highly charged and composed primarily of basic residues, but DNP packs against the more hydrophobic transmembrane (TM) helix 4 on the opposite side. The aromatic moiety of DNP contacts hydrophobic residue Ile183 and forms T-shaped aromatic stacking interactions with Tyr186. Moreover, one of the nitro groups forms electrostatic interactions with Arg234, whereas the hydroxyl group hydrogen bonds with Ser227. In this position, DNP occupies approximately the same space as the adenosine ring of ADP and the diterpene moiety of CATR (Extended Data Fig. 6a,b).

We subsequently docked FCCP, BAM15 and SF6487 into the c-state cavity, and found that they all bind in the pocket identified for DNP, with the best binding modes conserved between their neutral and negatively charged states (Fig. 4c and Extended Data Fig. 7a-d). Despite the chemical diversity of these compounds, their poses largely overlap such that key titratable atoms on all four molecules lie in the same spatial location (Extended Data Fig. 6e) with their aromatic rings against the hydrophobic side of the pocket.

We repeated the docking of DNP and other uncouplers using a homology model of the bovine transporter m-state based on a recent fungal AAC structure (PDB: 6GCI)⁴⁰. This revealed an interaction landscape for the molecules that largely resembled that observed in the c-state (Extended Data Fig. 7), supporting our experimental observation that cytosolic DNP activates I_H from both the c- and m-states of AAC (Extended Data Fig. 5b).

To validate this site, we performed molecular dynamics simulations of AAC1 in the c-state with charged DNP placed in solution away from the protein. Within 100 ns, DNP consistently entered the transporter driven by the strong positive potential in the cavity, and quickly moved to the location predicted from docking (Fig. 4b, Extended Data Fig. 8a,b and Supplementary Video 1). DNP in this site is dynamic, rapidly reorienting its hydroxyl and nitro substituents to interact with different protein moieties while largely remaining anchored to a single position on the protein through contact with its aromatic ring. We initiated additional simulations starting with DNP in the identified site either using neutral, protonated DNP or deprotonated DNP in a -160 mV membrane potential to bias negative charges toward the cytosol. In all simulations, DNP remained bound in the site for nearly the entire 500 ns duration (Extended Data Fig. 8b-d) and maintained interactions observed in the docking, as well as water-mediated contacts with Arg79, Arg187 and Asp231.

To determine whether FAs share any of the interactions predicted for chemical uncouplers, we initiated 14 molecular dynamics simulations of AAC1 in the c-state and 4 in the m-state with 16 FAs, either arachidonic or palmitic acid, placed randomly throughout the membrane (Supplementary Table 3). In 7 out of 14 c-state simulations, FAs from the outer leaflet entered the cavity through several distinct routes, while none bound to the m-state. In two trajectories, arachidonic acid initially occupies the chemical uncoupler site for over 1 μ s (Extended Data Fig. 9a,b and Supplementary Video 2). However, in all seven binding events, both arachidonic and palmitic acid eventually occupy a fenestration between TM5 and TM6 from which they do not leave. In many instances, the acyl chain protrudes into the lipid bilayer while the carboxylate interacts with residues Lys22, Arg79 and Arg279—all predicted to engage nucleotide phosphate groups^{38,39} (Fig. 4d and Extended Data Figs. 6f and 9c,d). The TM5–6 fenestration is evident in X-ray structures of the c-state in which it is occupied by several water molecules^{35,36}.

The configuration with the acyl tail anchored in the bilayer suggests a physical basis for our previous finding that a long hydrophobic tail is critical for FA activation of I_H through AAC and UCP1^{2,7}. Our data also reveal that both mitochondrial uncouplers and FAs bind within the water-accessible translocation pathway, but to different sites (Extended Data Fig. 6g). The uncoupler site overlaps with a portion of the CATR-binding site and the adenosine ring of docked ADP, whereas the region occupied by the dynamic FA headgroup overlaps with the carboxylate groups of CATR and the ADP phosphate groups (Extended Data Fig. 6a-d). We cannot exclude the possibility that FA exerts its effect from the uncoupler site rather than the TM5/6 fenestration site, as we observed a FA in both sites in the simulations (Extended Data Fig. 9a-d). Moreover, a simulation with two FAs placed in the cavity, one in each site, remains stable for a full microsecond (Extended Data Fig. 6h and Supplementary Video 3), showing that the c-state can simultaneously accommodate more than one FA molecule.

Biophysical model of H⁺ permeation through AAC

Continuum electrostatics calculations revealed a large, positive potential opposing H⁺ entry into the cytosolic cavity of the AAC1 c-state (Fig. 4e,f), consistent with its role as an anion transporter. Protons experience an electrostatic energy barrier (E_{elec}) of 10 $k_B T$ at the bottom of the cavity near the centre of the membrane ($z = 0$), where the protonatable groups of bound uncoupling agents are located (Fig. 4e and Extended Data Fig. 6e). Between $z = -5$ and -25 Å, there is no aqueous pathway through the closed gate on the matrix side and, therefore, another large energy barrier is present (Fig. 4f (grey region)). A negative membrane potential (normally present across the IMM) tilts the energy landscape in favour of H⁺ entry to the matrix, but does not significantly reduce the barrier (Fig. 4f and Extended Data Fig. 10a), consistent with a lack of I_H through AAC or UCP1 in the absence of FA^{2,7,19} or uncoupling agents (Fig. 1).

FAs and uncouplers probably enable H⁺ permeation by reducing the positive charge in the cavity and providing a protonatable group that acts as a stepping stone, similar to the FA–cofactor model proposed for UCP1⁶. However, overcoming the energetic barrier of the closed matrix gate requires a conformational change to open a H⁺ permeation pathway to the matrix. This pathway should be narrow to ensure selective H⁺ permeation, and

would therefore require only a minor conformational change short of the full c-state to m-state transition required for ADP/ATP exchange. We assume that both FAs and chemical uncouplers induce such a change as suggested by transient pre-steady-state currents recorded for AAC (and UCP1) in the presence of low- pK_a FA analogues after the application of a negative voltage^{2,7}.

On the basis of these assumptions, we created a mathematical model to examine the possibility of H^+ permeation through AAC once uncouplers bind and an H^+ -selective pathway to the matrix has opened. In this model, a proton first enters the transporter from the cytosol and binds to the uncoupler (pink molecule), followed by a transition to the narrow H^+ permeation pathway and eventual delivery to the matrix (Fig. 4g and Supplementary Note 1). The true nature of the H^+ pathway remains unclear as it may occur through a water wire, Brownian diffusion through a water filled passage, transfer to a protonatable site on the protein or another mechanism. Thus, we modelled this step as an Arrhenius rate, which can generally describe all of these processes at this coarse level.

We populated the rates in the model from the results of our structure-based computations on the c-state along with additional biophysical considerations (Supplementary Note 1). The model predicts a H^+ flux rate of around 15 H^+ per s at -160 mV in the presence of a DNP-like activator ($pK_a = 4.2$ in solution) and cytosolic and matrix pH values of 7 and 7.5, respectively (Fig. 4h; using the values shown in Supplementary Table 2). The large positive electrostatic barrier (E_{elec}) is a key determinant of the flux explaining the small unitary current. Even reducing E_{elec} from 10 $k_B T$ to 6 $k_B T$ causes an eightfold increase in I_H (Extended Data Fig. 10b). Thus, if a minor conformational change induced by the uncoupler/FA affects the shape of the cytosolic cavity and its electrostatic potential, it could significantly affect the H^+ transport rate. Furthermore, the same conformational change could position the uncoupler/FA deeper into the membrane electric field, which would increase the fraction of the membrane voltage (f) that H^+ move through as they bind from the cytosol. Increasing f from 0.1 (the computed value) to 0.5 causes a fourfold increase in the current (Extended Data Fig. 10c). Moreover, calculations with $f = 0.5$ better approximate the inward rectification observed in the experimental traces (Fig. 2 and Extended Data Fig. 2).

Tuning the pK_a of the uncoupler reveals that I_H is optimal above 5, and compounds with values under 2 do not induce current (Extended Data Fig. 10d), consistent with our earlier finding that low- pK_a FA analogues cannot induce I_H in either UCP1 or AAC^{2,7}. Uncouplers with moderately high pK_a values are still effective because the positive potential in the cavity essentially lowers their intrinsic pK_a making them more likely to give up their proton.

The largest structural uncertainty involves the nature of the H^+ pathway from the uncoupler to the matrix; however, the model places strong constraints on this rate (transition from state 2 to state 3 in Fig. 4g). The transition must occur in the upper nanosecond range or faster for efficient conduction or else H^+ unbinding to the cytoplasm dominates due to the large positive potential in the cavity (Extended Data Fig. 10e). Moreover, I_H is insensitive to the energy of the H^+ in the pathway provided that the forward rate from the uncoupler is fast,

consistent with a number of proton permeation mechanisms (further details are provided in Supplementary Note 1).

This model demonstrates the feasibility of the uncoupler-induced H^+ transport through the translocation pathway of AAC, replicates some of the measured electrophysiological properties and provides a molecular-based hypothesis that can drive future studies.

Discussion

Our results challenge the dogma that mitochondrial uncouplers such as DNP, FCCP, SF6847 and BAM15 induce I_H across the IMM through a protein-independent protonophoric mechanism⁴¹⁻⁴⁵. We demonstrate that the dominant mechanism of action of these uncouplers in the mitochondria is to induce I_H by activating AAC and UCP1, the same proteins that mediate the physiological, FA-induced mitochondrial I_H . However, at the level of the whole cell and the whole organism, the protonophoric activity of the uncouplers can still have a considerable impact due to their broad, indiscriminate effect on various cell membranes.

Electrophysiology and computational analysis suggest that chemical uncouplers and FAs induce I_H in AAC and UCP1 through the same general mechanism: both are weak acids that bind within the central translocation pathway to facilitate H^+ binding and permeation. Unexpectedly, FAs must be substantially more hydrophobic than chemical uncouplers to effectively activate I_H . Only long-chain FAs containing more than 12 carbons ($\log[P] > 4.6$) have the ability to activate AAC- and UCP1-dependent I_H in the low micromolar range^{2,7}. By contrast, DNP robustly activates I_H despite its much lower hydrophobicity (6 carbon atoms, $\log[P] = 1.67$). This indicates potential differences in the binding mechanisms of FAs and chemical uncouplers in the translocation pathways of AAC and UCP1, which are also confirmed by the molecular dynamics simulations and molecular docking (Supplementary Note 2).

All of the chemical uncouplers examined in this work have two mechanisms of action, a protonophoric activity affecting all cell membranes and an IMM-specific H^+ leak through AAC and UCP1. Despite this, our data indicate that it might be possible to develop specific activators of mitochondrial I_H through AAC (UCP1) that lack the protonophoric activity (Supplementary Note 1). Such next-generation anti-obesity and anti-diabetes drugs would activate native thermogenic pathways similar to their endogenous activators, long-chain FAs, and might have improved safety compared with general protonophores.

To date, UCP1 has been the primary pharmacological target to stimulate resting metabolic rate due to the thermogenesis that it induces in brown/beige fat. However, the impact of UCP1 in humans is limited because of a substantial reduction in brown adipose tissue with age and its near complete absence in certain individuals^{31,46}. AAC-dependent I_H can also increase energy expenditure by channelling the energy of mitochondrial substrate oxidation into heat. AAC is ubiquitously expressed and its contribution to overall body energy expenditure is therefore likely to exceed that of UCP1. By having two functions—ADP/ATP exchange and H^+ leak—AAC functions as a master regulator that controls the

partitioning of mitochondrial energy between ATP production and thermogenesis. Thus, this native, self-regulating pathway represents an attractive target for controlling energy expenditure.

Methods

Animals

Mice were maintained on a standard rodent chow diet under 12 h–12 h light–dark cycles. All of the animal experiments were performed with male mice according to procedures approved by the UCSF Institutional Animal Care and Use Committee. B6.129-Ucp1tm1Kz/J mice (abbreviated *UCP1*^{-/-}) were obtained from the Jackson Laboratory. The *AAC1*-deficient mice on the C57BL/6J2z background were obtained from the laboratory of D. Wallace²⁴. WT and *UCPT*^{-/-} mice (B6.129-Ucp1tm1Kz/J) on the C56BL/6J background were obtained from the Jackson Laboratory. For the experiments with *AAC1*^{-/-} mice, we used WT C57BL/6J2z and C56BL/6J control mice with the same results. For all of the other knockout mouse strains C56BL/6J mice were used as control. All mice were aged 2 months to 1 year.

C2C12 cell culture and generation of *AAC1/AAC2* knockout cells

The original WT C2C12 cell line (immortalized mouse myoblast cell line) was purchased from ATCC (<https://www.atcc.org/>) and cultured in complete Dulbecco's modified Eagle's medium (DMEM; DMEM with 25 mM glucose, 10% fetal bovine serum (FBS), penicillin and streptomycin antibiotics). The C2C12 cell line was authenticated by ATCC. The *AAC1/AAC2* DKO C2C12 cells were generated by Alstem (<http://www.alstembio.com/>) using the CRISPR–Cas9 system⁷. Alstem provided initial authentication of *AAC1/AAC2* DKO C2C12 cells using PCR. We provided further authentication of the DKO cells with western blotting. All cell lines were tested negative for mycoplasma contamination. Transfection of gRNA *Cas9* plasmids was performed using the Invitrogen Neon transfection system. gRNA candidates were selected using PCR amplification and selected for single-cell clone isolation.

Isolation of mitochondria and mitoplasts from tissues

Mice were euthanized by CO₂ asphyxiation followed by cervical dislocation. For the preparation of mitoplasts from heart, skeletal muscle, brown fat, kidneys and liver, the selected mouse tissues were isolated, rinsed and homogenized in ice-cold medium containing 250 mM sucrose, 10 mM HEPES, 1 mM EGTA and 0.1% bovine serum albumin (BSA) (pH adjusted to 7.2 with Trizma base) using a glass grinder with six slow strokes of a Teflon pestle rotating at 275 (soft tissues) or 600 (fibrous tissues) rotations per minute. The homogenate was centrifuged at 700g for 5–10 min to pellet nuclei and unbroken cells. For some tissues, the first nuclear pellet was resuspended in the same solution and homogenized again to increase the yield of mitochondria. Mitochondria were collected by centrifugation of the supernatant at 8,500g for 10 min.

Mitoplasts were produced from mitochondria using a French press. In brief, mitochondria were suspended in a solution containing 140 mM sucrose, 440 mM D-mannitol, 5 mM

HEPES and 1 mM EGTA (pH adjusted to 7.2 with Trizma base) and then processed using a French press at 1,200–2,000 psi to rupture the outer membrane. Mitoplasts were pelleted at 10,500g for 15 min and resuspended for storage in 500 μ l of solution containing 750 mM KCl, 100 mM HEPES and 1 mM EGTA (pH adjusted to 7.2 with Trizma base). Mitochondria and mitoplasts were prepared at 0–4 °C and stored on ice for up to 5 h. Immediately before the electrophysiological experiments, 15–50 μ l of the mitoplast suspension was added to 500 μ l solution containing 150 mM KCl, 10 mM HEPES and 1 mM EGTA (pH adjusted to 7.0 with Trizma base) and plated on 5 mm coverslips pretreated with 0.1% gelatin to reduce mitoplast adhesion.

Patch-clamp recordings

Whole-IMM patch-clamp recording was performed from isolated mitoplasts^{2,7,19}. The mitoplasts used for patch-clamp experiments were 3–5 μ m in diameter and typically had membrane capacitances of 0.3–1.2 pF. Gigaohm seals were formed in the bath solution containing 150 mM KCl, 10 mM HEPES and 1 mM EGTA (pH 7 adjusted using Trizma base). Voltage steps of 250–500 mV and 1–50 ms were applied to break-in into the mitoplast and obtain the whole-mitoplast configuration, as monitored by the appearance of capacitance transients. Mitoplasts were stimulated every 5 s. Currents were normalized to membrane capacitance to obtain current densities (pA pF⁻¹).

All indicated voltages are on the matrix side of the IMM (pipette solution) as compared to the cytosolic side (bath solution, defined to be 0 mV). Normally, currents were induced by a voltage ramp from –160 mV to +100 mV to cover all physiological voltages across the IMM. In the figures, voltage protocols are shown above the current traces. Currents flowing into mitochondria are shown as negative, whereas those flowing out are positive. Membrane capacitance transients observed after the application of voltage steps were removed from current traces.

Both the bath and pipette solutions were formulated to record H⁺ currents and contained only salts that dissociate into large anions and cations normally impermeant through ion channels or transporters. The pipettes were filled with 130 mM tetramethylammonium hydroxide, 1.5 mM EGTA, 2 mM Tris chloride and 100 mM HEPES. pH was adjusted to 7.5 with D-gluconic acid, and tonicity was adjusted to around 360 mmol kg⁻¹ with sucrose. Typically, pipettes had resistances of 25–35 M Ω . and the access resistance was 40–75 M Ω . Whole-mitoplast I_H was recorded in the bath solution containing 100 mM HEPES and 1 mM EGTA (pH 7.0 with Trizma base, and tonicity ~300 mmol kg⁻¹ with sucrose). Whole-mitoplast K⁺ current activated by valinomycin was recorded in the bath solution containing 50 mM KOH, 182 mM HEPES and 1 mM EGTA (pH 7.0 with HEPES, tonicity 300 mmol kg⁻¹).

In the absence of significant cytosolic triglyceride storage in non-adipose tissues, membrane phospholipid hydrolysis may be an important source of free FA for I_H activation in vivo. To record I_H through UCP1, before the application of uncouplers, the endogenous membrane FA were removed by a 30–40 s pretreatment with 10 mM M β CD.

In the experiments that compared I_H activated by protonophores in IMM versus plasma membrane, whole-IMM measurements were performed from skeletal muscle mitoplasts as described above. Whole-cell plasma membrane H^+ current measurements were performed from HEK293 cells. Gigaohm seals with HEK293 cells were formed in a Ringer solution containing 150 mM NaCl, 4 mM KCl, 2 mM $CaCl_2$, 1 mM $MgCl_2$, 5.6 mM glucose and 25 mM HEPES (pH 7.4 with NaOH). Whole-IMM and whole-cell I_H measurements were performed under identical recording conditions. A large pH gradient was set up between the bath solution (pH 6.0) and pipette solution (pH 7.5) to facilitate the detection of small H^+ currents. Currents were expressed as current densities ($pA\ pF^{-1}$) to enable comparison between the small mitoplasts and much larger plasma membranes. The pipettes were filled with 130 mM tetramethylammonium hydroxide, 1.5 mM EGTA, 2 mM Tris chloride and 100 mM HEPES (pH 7.5 with D-gluconic acid, tonicity of around 360 mmol kg^{-1} with sucrose). The bath solution contained 100 mM MES and 1 mM EGTA (pH 6.0 with Trizma base, and tonicity of around 300 mmol kg^{-1} with sucrose).

All of the experiments were performed under continuous perfusion of the bath solution. All electrophysiological data presented were acquired at 10 kHz and filtered at 1 kHz.

C2C12 mitochondrial isolation

C2C12 cells (WT or DKO) from 10 15 cm dishes were trypsinized and collected with ice-cold PBS at 1,000g for 5 min. The cell pellet was incubated with hypotonic buffer (20 mM HEPES, pH 7.5, 5 mM KCl, 1.5 mM $MgCl_2$ and 1 mg ml^{-1} essentially FA-free BSA) on ice for 10 min. Cells were then homogenized (25 strokes) using a tight-fitting glass dounce tissue grinder and rapidly made isotonic by adding the homogenate to 2/3 volume of 2.5 \times MSH (20 mM HEPES, pH 7.5, 525 mM mannitol, 175 mM sucrose, 5 mM EDTA, and 1 mg ml^{-1} essentially FA-free BSA). The homogenate was centrifuged at 600g for 10 min, and the supernatant was then centrifuged at 8,500g for 10 min to pellet mitochondria. Mitochondria were resuspended in MSH buffer (20 mM HEPES, pH 7.5, 210 mM mannitol, 70 mM sucrose and 2 mM EDTA). Mitochondrial protein was quantified using the bicinchoninic acid assay (Thermo Fisher Scientific).

Mitochondrial respiration using the Seahorse Extracellular Flux Analyzer

In a volume of 50 μ l, 30 μ g of mitochondrial protein was added to each well of a XF cell culture microplate (on ice). The mitochondria-loaded XF microplate was centrifuged at 2,000g for 20 min for adherence of organelles. Respiration buffer (20 mM Tris, pH 7.4, 210 mM mannitol, 70 mM sucrose, 0.1 mM EGTA, 3 mM $MgCl_2$, 5 mM KH_2PO_4 , 0.1% essentially FA-free BSA, 10 mM sodium pyruvate and 5 mM malate) was added to a final volume of 0.5 ml, and the XF microplate was warmed at 37 $^{\circ}C$ for 8 min before loading into a XFe24 Extracellular Flux Analyzer (Seahorse Bioscience). The OCRs were recorded using a mix (20 s) and measure (2 min) cycle.

Mitochondrial respiration using a Clark electrode

Mitochondrial respiration was assessed in a 0.4 ml continuously stirred chamber containing the standard incubation medium, containing 125 mM KCl, 0.5 mM $MgCl_2$, 3 mM KH_2PO_4 , 10 mM HEPES, pH 7.4, 10 μ M EGTA and supplemented with 3 mM pyruvate plus 1 mM

malate. The chamber was maintained at 37 °C and was equipped with a custom-made Clark-type oxygen electrode and a tightly sealed lid. The slope of the oxygen consumption trace corresponded to the respiratory rate. Linear regression was performed on sections of the respiration traces for the 60 s immediately preceding and immediately after the addition of DNP to the mitochondria. The linear equation yielded during each linear regression analysis was used to determine the change in volts per minute for the trace segment of interest. Using the total known scale in volts for each trace and an experimentally determined value, in nmol, for the amount of oxygen dissolved in an aqueous solution at 37 °C and 1 atm, electrical potential was converted to nmol O₂. The known amount of dissolved O₂ under normal air pressure in 0.15 M KCl solution at 37 °C has previously been determined to be 162.4 nmol O₂ per ml (ref.⁴⁷). Finally, the amount of nmol O₂ consumed per minute was normalized to mitochondrial protein (275 µg mitochondria/400 µl volume of the chamber).

Cell respiration analysis using the Oroboros O₂k respirometer

C2C12 cell culture was maintained in DMEM supplemented with 25 mM glucose, 1 mM sodium pyruvate, 4 mM GlutaMAX and 10% FBS. The respiration medium was DMEM (containing 5.6 mM glucose, 1 mM sodium pyruvate, 4 mM L-glutamine, 20 mM HEPES, 10% FBS and no phenol red) with approximately 325 mOsm, at around pH 7.15. The cell concentration in each 2 ml respiration chamber was 5×10^6 cells per ml. The temperature was maintained at 37 °C and the cell suspension was stirred continuously at 750 rpm. After the oxygen consumption rate reached steady-state, 1 µM oligomycin was first applied to inhibit ATP synthase. Increasing concentrations of uncouplers (FCCP, 2,4-DNP and BAM15) or valinomycin were then added as indicated. Nigericin (K⁺/H⁺ antiporter, 1 µM) was added to reduce matrix K⁺ accumulation and mitochondrial swelling caused by valinomycin²⁷. At the end of the experiments, 1 µM rotenone and 2.5 µM antimycin were applied to inhibit mitochondrial respiration and determine the non-mitochondrial respiration.

Immunoblot analysis

For western blot analysis, cells were lysed in RIPA buffer (1% igepal, 0.1% sodium dodecyl sulfate, 0.5% sodium deoxycholate, 150 mM NaCl, 1 mM EDTA, 50 mM Tris-HCl (pH 7.4) and a cocktail of protease inhibitors). Lysates were resolved by SDS-PAGE, transferred to PVDF membrane (Millipore), and probed with anti-Na⁺/K⁺-ATPase (Abcam, ab76020, GR3237646-13, 1:10,000), anti-TOMM20 (Sigma Prestige, HPA011562, 000012083, 1:500), anti-AAC1 (Abcam, ab110322, R4759, 1:500) and anti-AAC2 (Cell Signaling Technology, CST 14671, 1:500) antibodies and OXPHOS antibody cocktail for mitochondrial respiratory content (Abcam, ab110413, MS604/Q5039, 1:500, NDUFB8 (complex I), SDHA (complex II), core 2 subunit (complex III), CIV-I subunit (complex IV) and ATP5A (complex V)).

Cell lines

The C2C12 WT cell line was purchased from ATCC (<https://www.atcc.org/>). These WT C2C12 cells were used to generate the double AAC1/AAC2 KO C2C12 (DKO) cell line. The DKO cell line was generated by Alstem (<http://www.alstembio.com/>) using the Crispr-Cas9 system⁷. The C2C12 cell line was authenticated by ATCC. The AAC1/AAC2 double KO C2C12 (DKO) cell line was generated and first authenticated by Alstem. We provided

further authentication of the DKO cells with western blotting. All cell lines tested negative for mycoplasma contamination. The HEK293T cells (*Homo sapiens*) were provided and authenticated by the UCSF Cell Culture Facility (<https://cgec.ucsf.edu/cell-culture-and-banking-services>).

Chemicals

The following chemicals were used in this study: carboxyatractyloside potassium salt (CATR, Sigma-Aldrich, 4992), bongkreikic acid triammonium salt (BKA, Millipore 203671), guanosine 5'-diphosphate tris salt (GDP, Sigma-Aldrich, G7252), adenosine 5'-diphosphate sodium salt (ADP, Sigma-Aldrich, A2754), adenosine 5'-triphosphate disodium salt hydrate (ATP, Sigma-Aldrich, A6419), methyl- β -cyclodextrin (M β CD, Sigma-Aldrich, C4555), BAM15 (Sigma-Aldrich, SML1760) and SF6847 (CAS10537-47-0, Calbiochem, Sigma-Aldrich).

Statistical analysis

Data are presented as mean \pm s.e.m. as specified in the figure legend. Statistical analysis was performed using GraphPad Prism 8. Statistical significance with exact *P* values was determined using the method indicated in the corresponding figure legend.

Docking methods

The Small Molecule Drug Discovery Suite 20191 (Schrödinger) was used for all docking calculations. The protein was prepared in a ready-to-dock-format with the Protein Preparation Wizard workflow. For docking to the c-state, PDB 2C3E (ref. ³⁶) was used, whereas docking to the m-state used the homology model prepared for molecular dynamics simulations. Protein hydroxyl group orientations and protonation states were assigned at pH 7 using PROPKA⁴⁸ in Epik v.4.7. Small molecules were drawn using Maestro v.11.9, and their protonation states and tautomers were assigned at pH 7.0 ± 3 in Epik v.4.7. Geometry was optimized and partial charges were assigned using Jaguar v.10.3 using the B3LYP-D3 density functional⁴⁹ to account for non-covalent dispersion interactions with the 6-31G** and 6-31**+ basis sets for neutral states and anions, respectively, with ultra-fine level of accuracy of the calculation grid. Other parameters for proteins and small molecules used in the docking experiments came from the OPLS3e force-field⁵⁰. A $30 \times 32 \times 30$ Å docking grid was erected in the central cavity, and molecules docked in flexible mode allowing bond rotation while enforcing aromatic planarity restricting amide groups to *trans*, treating halogens as hydrogen-bond acceptors, and aromatic hydrogens as donors. For docking into the c-state, the protein was held rigid and the Glide XP⁵¹ scoring function was used, including strain energy for the ligands in the final score and returning a maximum of five optimal poses. For docking into the m-state, the induced fit workflow was applied, using the alanine trimming procedure along with the Glide SP scoring function for rescoring, and a maximum of 20 poses were returned. Final poses were picked on the basis of a combination of visual inspection, chemical intuition and score.

Molecular dynamics simulations

Simulations of the c-state of AAC1 were prepared using PDB 2C3E (ref. ³⁶), while simulations of the m-state were prepared using a homology model based on PDB 6GCI (ref. ⁴⁰), with the final simulation model in both cases consisting of residues 1–297 of bovine AAC1 and 3 bound cardiolipin lipids. Homology modelling of the bovine AAC1 m-state was performed using the automated SWISS-MODEL web server⁵², using the BKA-bound *Thermothelomyces thermophilus* AAC m-state structure (PDB: 6GCI) as a template (48% sequence identity to bovine AAC1). The inhibitors CATR and BKA found in the crystal structures were removed for all of the simulations. Four missing residues at the protein C terminus in PDB 2C3E were added manually in PyMOL (The PyMOL Molecular Graphics System, v.1.8 Schrödinger) as an extension of TM6. For both states, three tetra-linoleoyl cardiolipin lipids were built into lipid-binding sites manually in PyMOL using the partial lipid residues found in the crystal structures as templates. Structures were embedded in lipid bilayers using the membrane builder application of the CHARMM-GUI web server⁵³. Some bilayers were built using POPC as the primary lipid, while others used a 1:1 mixture of PC:PE lipids with additional cardiolipin molecules to more closely mimic the composition of the mitochondrial inner membrane⁵⁴ (Supplementary Table 3). The lipid composition of the bilayer had no apparent effect on the structure or ligand-binding properties of AAC1. For simulations using FA in the bilayer, FA molecules were also placed randomly using the CHARMM-GUI membrane builder. For simulations using DNP, the DNP molecule was placed either randomly in the aqueous portion of the simulation box, or in the AAC1-binding site on the basis of previous simulation runs. The force field parameters used for protein and lipid/FA were CHARMM36m⁵⁵ and CHARMM36⁵⁶, respectively. Water was represented with the TIP3P model⁵⁷, and standard CHARMM ion parameters were used⁵⁸. CHARMM-compatible parameters for neutral and deprotonated DNP were obtained for CgenFF (v.4.0)⁵⁹ using the paramchem web server (<https://cgenff.umaryland.edu/>), and atomic partial charges were reoptimized using the Force Field Toolkit VMD plugin⁶⁰ and gaussian09 (ref. ⁶¹). Additional simulation details are provided in Supplementary Table 3.

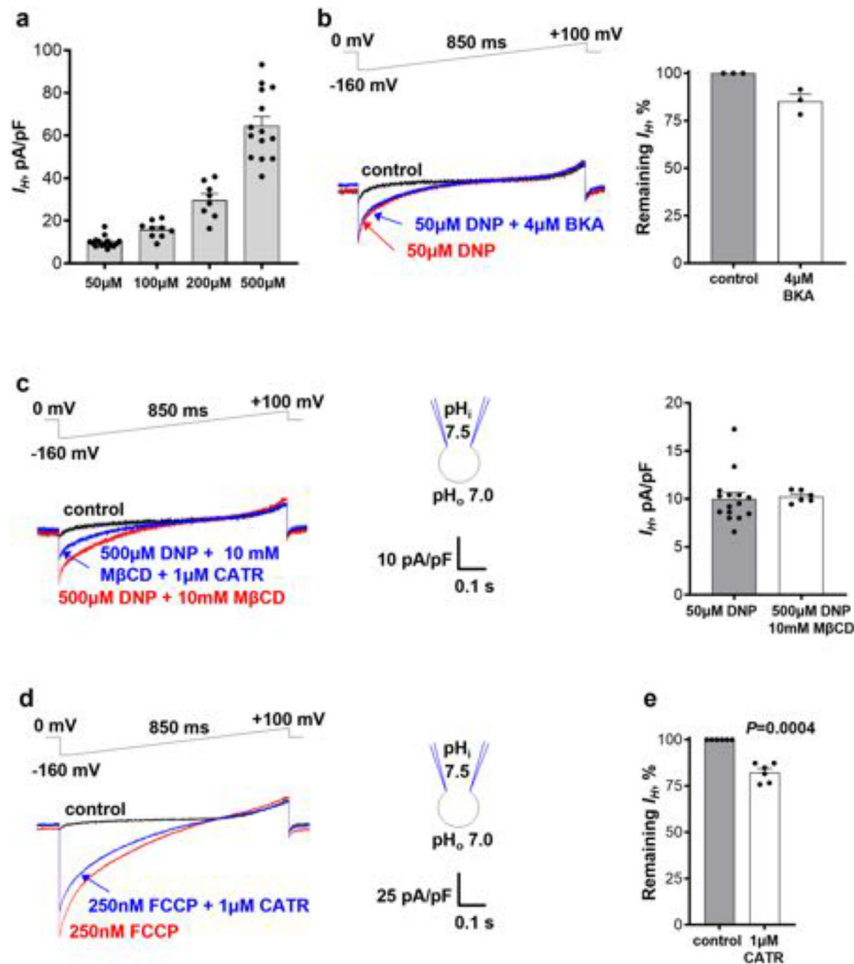
All molecular dynamics simulations were performed on local GPU computing resources using GROMACS (v.2018 or v.2020)⁶². Systems were initially minimized with 2,000 kJ mol⁻¹ nm² harmonic restraints on all protein heavy atoms and DNP or FA heavy atoms, when applicable, followed by a multistep equilibration in which protein restraints were gradually reduced over 10 ns. Production simulations were run using a 2 fs timestep, a Nose–Hoover (GROMACS v.2018) or stochastic-velocity rescaling (GROMACS v.2020) thermostat at 315 K (refs. ^{63–65}), and a semi-isotropic Parrinello–Rahman barostat with a pressure of 1 atm (ref. ⁶⁶). A 12 Å cut-off was used for Lennard–Jones interactions with force switching in the range of 10–12 Å, and long range electrostatics were calculated using the particle mesh Ewald method⁶⁷. Bonds to hydrogen atoms were constrained using the LINCS algorithm⁶⁸. Atomic coordinates were saved every 250 ps for all simulations. For systems simulated with an applied membrane potential, we used the constant electric field method where $E_{\text{applied}} = V/L_z$ where L_z is the length of the simulation box in the z dimension⁶⁹, and the membrane potential was equilibrated at a rate of 10 mV per 5 ns. Simulation analysis was performed using the MDAnalysis Python package⁷⁰. Molecular

graphics were generated using the Visual Molecular Dynamics (VMD)⁷¹ and PyMOL software packages.

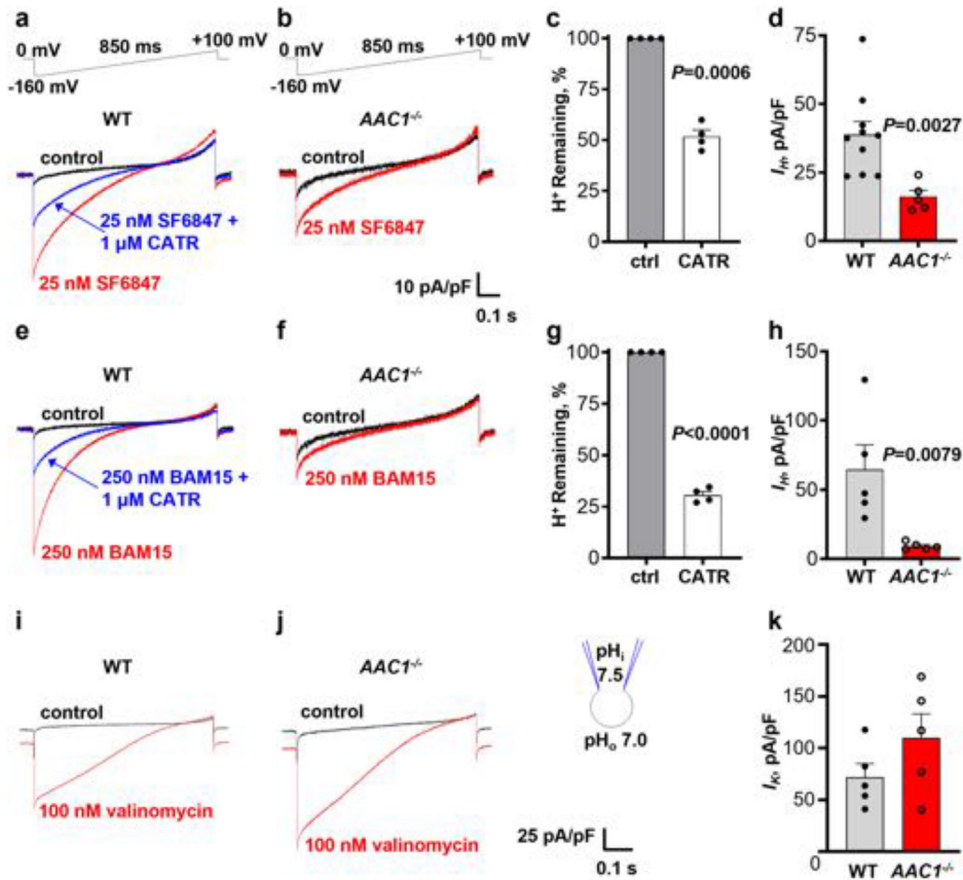
Electrostatic calculations

Continuum electrostatics calculations were carried out on a representative snapshot of bovine AAC1 extracted from a molecular dynamics simulation at 75 ns (ID 2 in Supplementary Table 3). The protein only was embedded in a membrane-like environment using APBSmem⁷² and calculations were performed using APBS⁷³ with the PARSE parameter set for membrane voltage calculations and Swanson for ion solvation energy calculations (a list of additional parameters is provided in Supplementary Table 1; see ref. ⁷² for a more complete discussion of the electrostatic parameters, including the rationale for the standard dielectric values for the protein, membrane core and membrane headgroups used here). Ion solvation energies, accounting for all fixed and mobile charge interactions and the ion self-energy, were computed as described in ref.⁷² using nonlinear Poisson–Boltzmann theory (npbe), and the influence of the membrane potential was determined from separate calculations using the linearized form of the theory with appropriate boundary conditions (lpbe). Both calculations were added together to produce the composite curves shown in Fig. 4f.

Extended Data

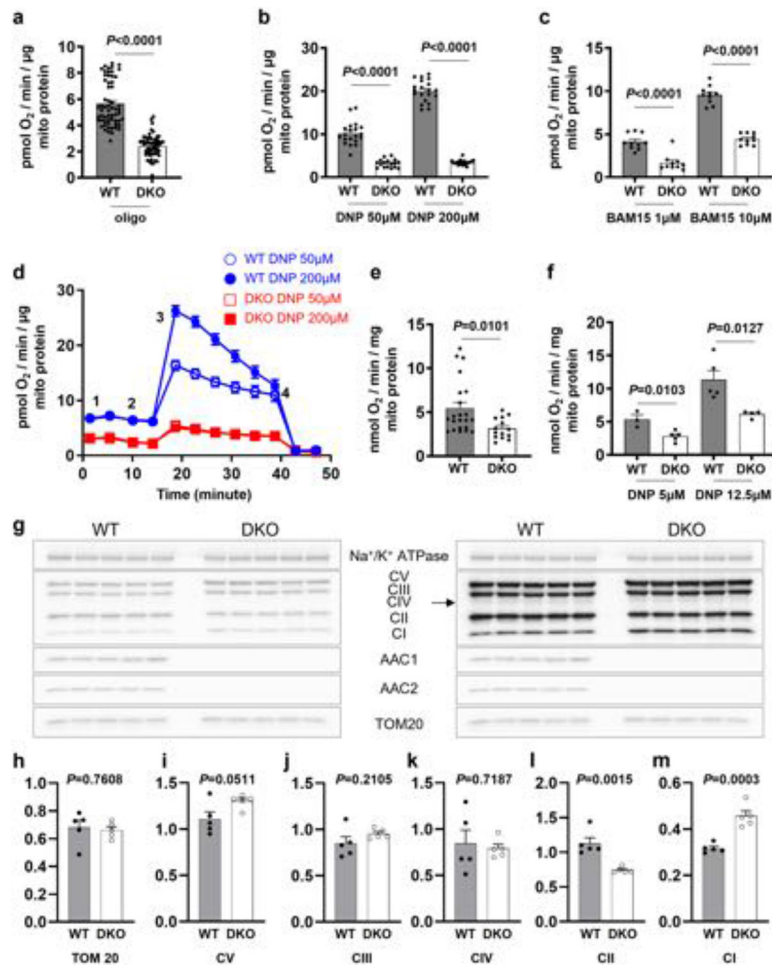
**Extended Data Fig. 1 l. DNP- and FCCP-induced mitochondrial I_H .**

a, I_H densities induced by different DNP concentrations in heart IMM at -160 mV ($n = 15$ for $50 \mu\text{M}$, $n = 9$ for $100 \mu\text{M}$, $n = 8$ for $200 \mu\text{M}$, $n = 14$ for $500 \mu\text{M}$). Data are mean \pm SEM. **b**, I_H induced by $50 \mu\text{M}$ DNP in heart IMM before and after application $4 \mu\text{M}$ BKA (left). Remaining fraction of I_H induced by $50 \mu\text{M}$ DNP at -160 mV after inhibition by $4 \mu\text{M}$ BKA, $n = 3$ (right). Data are mean \pm SEM. **c**, I_H induced by $500 \mu\text{M}$ DNP on the background of 10 mM M β CD in heart IMM before and after application of $1 \mu\text{M}$ CATR (left). I_H densities induced by $50 \mu\text{M}$ DNP alone ($n = 15$) and $500 \mu\text{M}$ DNP on the background of 10 mM M β CD ($n = 6$) (right). Data are mean \pm SEM. **d**, I_H induced by 250 nM FCCP in heart IMM was inhibited by $1 \mu\text{M}$ CATR. **e**, Remaining fraction of FCCP-induced I_H at -160 mV after inhibition by $1 \mu\text{M}$ CATR. Control (100% I_H) is shown in gray. Data are mean \pm SEM. Paired t-test, two-tailed, control vs. $1 \mu\text{M}$ CATR, $n = 6$.



Extended Data Fig. 2 l. SF6847 and BAM15 induce IH via AAC.

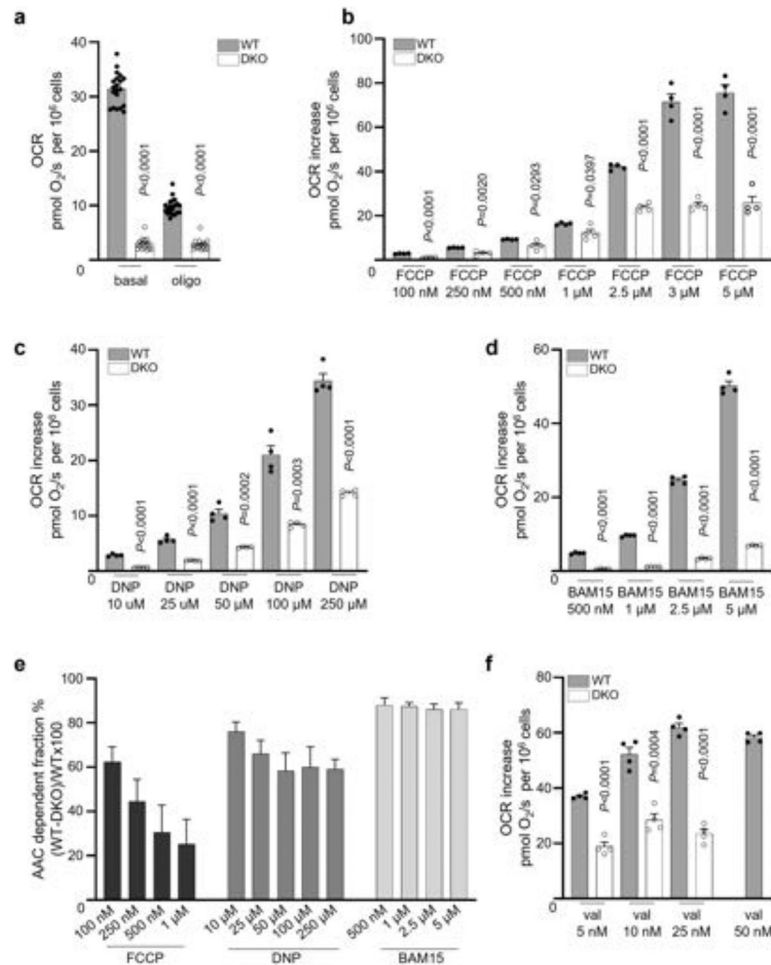
a, I_H induced by 25 nM SF6847 across WT heart IMM before and after application 1 μM CATR. **b**, I_H induced by 25 nM SF6847 in *AAC1*^{-/-} heart IMM. **c**, Remaining fraction of I_H induced by 25 nM SF6847 at -160 mV after inhibition by 1 μM CATR. Data are mean ± SEM. Paired t-test, two-tailed, control vs CATR, *n* = 6. **d**, I_H densities induced by 25 nM SF6847 at -160 mV in WT (*n* = 10) and *AAC1*^{-/-} (*n* = 5) heart IMM. I_H was measured at -160 mV. Data are mean ± SEM. Mann-Whitney U-test, two-tailed, WT vs *AAC1*^{-/-}. **e**, I_H induced by 250 nM BAM15 across WT heart IMM before and after application 1 μM CATR. **f**, I_H induced by 250 nM BAM15 in *AAC1*^{-/-} heart IMM. **g**, Remaining fraction of I_H induced by 250 nM BAM15 at -160 mV after inhibition by 1 μM CATR. Data are mean ± SEM. Paired t-test, two-tailed, control vs CATR, *n* = 4. **h**, I_H densities induced by 250 nM BAM15 at -160 mV in WT (*n* = 5) and *AAC1*^{-/-} (*n* = 5) heart IMM. I_H was measured at -160 mV. Data are mean ± SEM. Mann-Whitney U-test, two-tailed, WT vs *AAC1*^{-/-}. **i**, **j**, Representative K⁺ currents induced by 100 nM valinomycin across heart IMM in WT (**i**) and *AAC1*^{-/-} (**j**). **k**, K⁺ current densities induced by 100 nM valinomycin at -160 mV in WT and *AAC1*^{-/-} heart IMM (*n* = 5). Data are mean ± SEM.



Extended Data Fig. 3 l. DNP- and BAM-induced mitochondrial uncoupled respiration require AAC.

a. Oxygen consumption rate (OCR) of mitochondria isolated from WT and DKO C2C12 cells in the presence of oligomycin as measured with the Seahorse Analyzer. Each point corresponds to an individual respiration well, $n = 60$ (WT), and $n = 56$ (DKO). Mann–Whitney U -test, two-tailed. Data represent mean \pm SEM. **b, c** OCR increase observed in mitochondria isolated from WT and DKO C2C12 cells after addition of DNP, $n = 20$, WT and $n = 17$, DKO (**b**) or BAM15, $n = 10$, WT and $n = 10$, DKO (**c**) in the presence of oligomycin. Each point corresponds to OCR change in an individual well of the Seahorse Analyzer, $n = 17 - 20$ (DNP) and $n = 10$ (BAM15). Data are mean \pm SEM. Mann–Whitney U -test, two-tailed. **d.** OCR time course of isolated mitochondria from WT ($n = 20$ wells, blue) and DKO C2C12 cells ($n = 17$ wells, red). Basal respiration (1), addition of oligomycin (2), addition of DNP (3) and addition of rotenone/antimycin (4). Data represent mean \pm SEM. **e.** Basal OCR of mitochondria isolated from WT or DKO C2C12 cells measured with a Clark electrode ($n = 22$, WT and $n = 14$, DKO). Mann–Whitney U -test, two-tailed. Data represent mean \pm SEM. **f.** OCR increase above basal, observed using a Clark electrode after addition of DNP to mitochondria isolated from WT and DKO C2C12 cells; $n = 3$ (WT) and $n = 5$ (DKO) for $5 \mu\text{M}$; $n = 5$ (WT) and $n = 4$ (DKO) for $12.5 \mu\text{M}$. Data are mean \pm SEM. Mann–Whitney U -test, two-tailed. **g, Left**

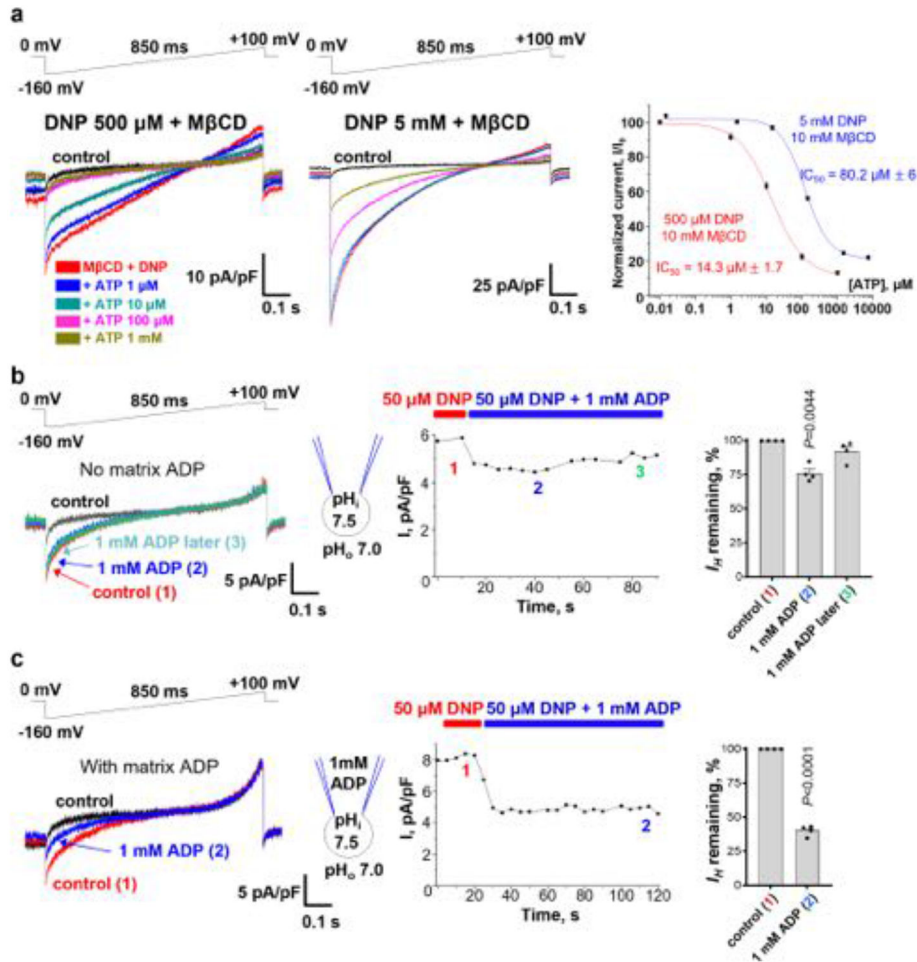
panel: Immunoblots of WT ($n = 5$) and DKO ($n = 5$) C2C12 cells showing expression of the electron-transport chain complexes complex I (CI) – complex V (CV), TOM20, and the loading control (plasma membrane Na^+/K^+ ATPase). *Right panel:* The same immunoblots at longer exposure for the bands corresponding to complexes I-V. **h**, TOM20 protein expression level relative to Na^+/K^+ ATPase for the immunoblot shown in **(g)**. Data shown as mean \pm SEM, $n = 5$. Two-tailed unpaired t-test, WT vs DKO. **i–m**, Protein expression levels for complexes I-V relative to TOM20 for the immunoblots shown in **(g)**. Data shown as mean \pm SEM, $n = 5$. Two-tailed unpaired t-test, WT vs DKO. For gel source data, see Supplementary Fig. 1.



Extended Data Fig. 4 l. Respiration induced by mitochondrial uncouplers in intact cells depends on AAC.

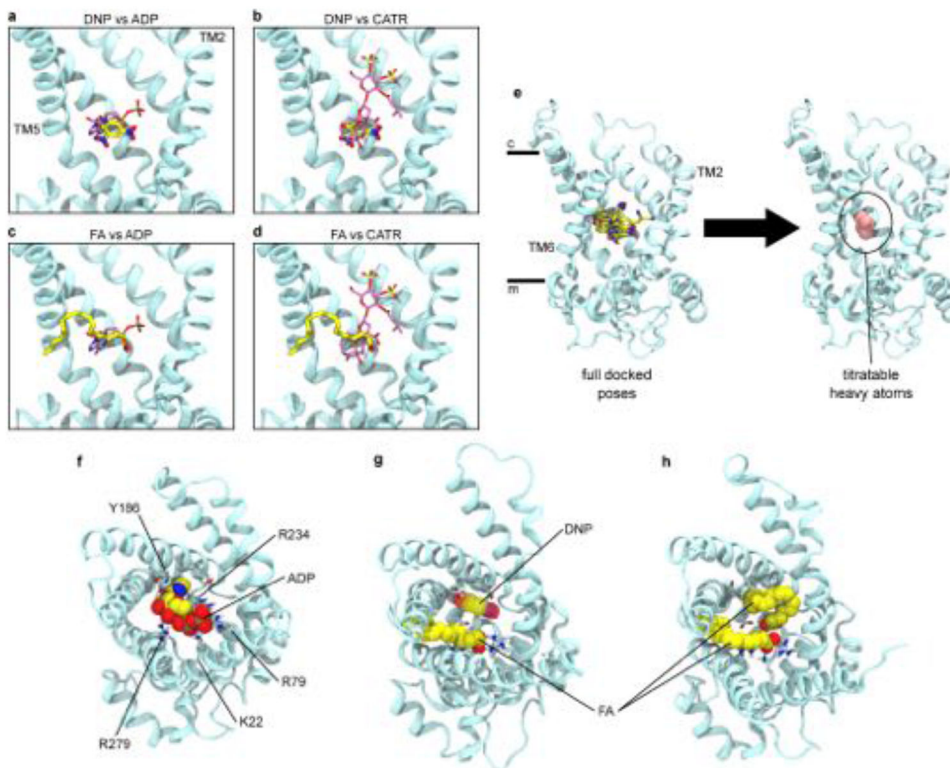
a, Oxygen consumption rate (OCR) of WT ($n = 20$) and DKO C2C12 cells ($n = 16$) before and after addition of oligomycin (oligo). Two-tailed unpaired t-test, WT vs DKO. **b–d**, OCR increase observed in WT and DKO C2C12 cells upon application of various concentrations of FCCP ($n = 4$), DNP ($n = 4$), and BAM15 ($n = 4$). All OCRs were measured in the presence of oligomycin. Two-tailed unpaired t-test, WT vs DKO. **e**, AAC-dependent fraction of uncoupled respiration at various concentrations of FCCP, DNP, and BAM15. All data shown as mean \pm SEM, $n = 4$. **f**, OCR increase observed in WT and DKO C2C12 cells

upon application of various concentrations of valinomycin. All OCRs were measured in the presence of oligomycin. Nigericin (1 μM) was added to reduce matrix K^+ accumulation and mitochondrial swelling caused by valinomycin²⁷. Two-tailed unpaired t-test, WT vs DKO, $n = 4$.



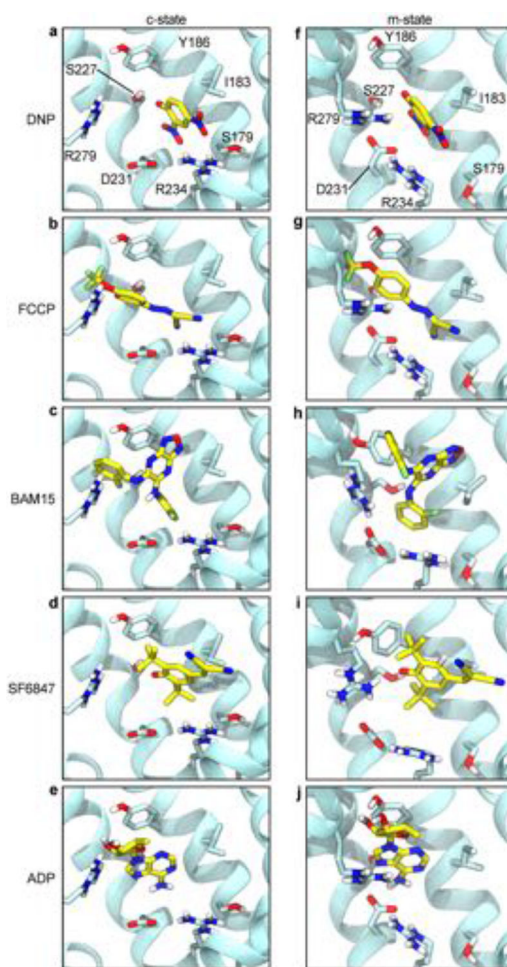
Extended Data Fig. 5 l. Adenine nucleotides inhibit DNP-induced I_H .

a, DNP-induced I_H at various ATP concentrations on the cytosolic face of the IMM of brown fat. I_H was activated with either 500 μM DNP (left panel) or 5 mM DNP (middle panel) applied in the presence of 10 mM M β CD. Control currents were recorded in 10 mM M β CD. Right panel is the dose-dependence of I_H inhibition by ATP at 500 μM DNP (red, $n = 5$) and 5 mM DNP (blue, $n = 6$). Current amplitudes were measured at -160 mV. Data are mean \pm SEM. **b**, Left panel, I_H induced by 50 μM DNP in control (1), upon transient inhibition by bath ADP (2), and upon subsequent recovery in the presence ADP (3). Heart mitoplast. Middle panel, DNP-dependent I_H time course of left panel. I_H was measured at -160 mV. Right panel, ADP inhibition of I_H at points (2) and (3) in left and middle panels. Data are mean \pm SEM. Paired t-test, two-tailed, ADP (2) vs. control, $n = 4$. **c**, I_H activated by 50 μM DNP was inhibited by bath ADP. Pipette solution contained 1 mM ADP. Heart mitoplast. Middle panel, I_H time course of the left panel. Right panel, ADP inhibition of I_H in point (2) as in left and middle panels. Data are mean \pm SEM. Paired t-test, two-tailed, $n = 4$.



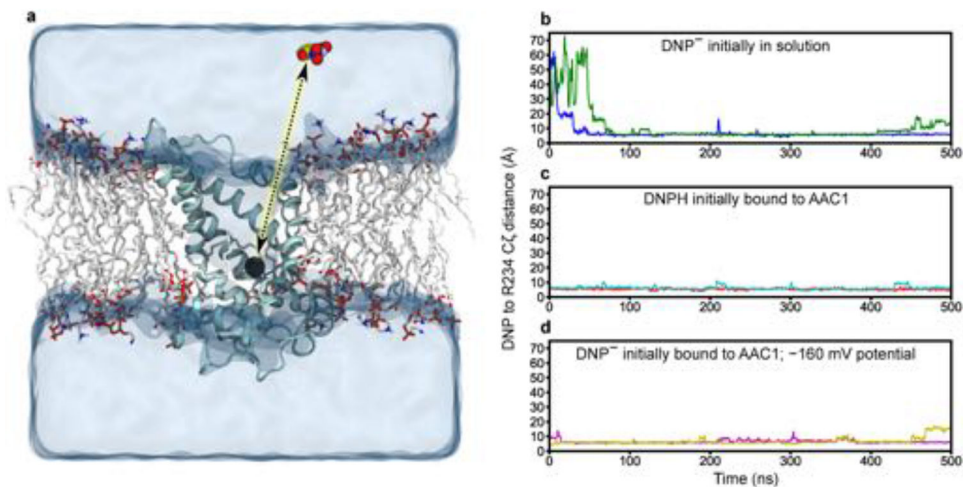
Extended Data Fig. 6 l. Binding mode comparisons of CATR, ADP, small molecule uncouplers, and fatty acid.

a–d, Comparison of binding poses in the AAC1 c-state (2C3E) cavity of **(a)** docked ADP and docked DNP; **(b)** crystallographic CATR and docked DNP; **(c)** FA (arachidonic acid) pose from MD simulation and docked ADP; and **(d)** FA pose from MD simulation and crystallographic CATR. TM helices 1 and 6 of AAC1 are hidden. ADP and CATR are shown as thinner sticks with mauve carbon atoms. DNP and FA are shown as thicker sticks with yellow carbon atoms. **e**, Full final docked DNP, FCCP, BAM15, and SF6847 poses in their deprotonated and protonated forms (left) as in Fig. 4c and titratable heavy atoms (pink spheres) from these poses (right). TM helix 1 of AAC1 is hidden. **f**, ADP binding mode to the c-state as predicted from docking. **g**, Simulation snapshot showing arachidonic acid (solid spheres) bound to the TM5/6 fenestration of AAC1 in the c-state superimposed with the docked pose of DNP (transparent spheres). **h**, Simulation snapshot showing two arachidonic acid molecules bound simultaneously to the AAC1 c-state cavity, one in the TM5/6 fenestration and the other in the small molecule uncoupler site.



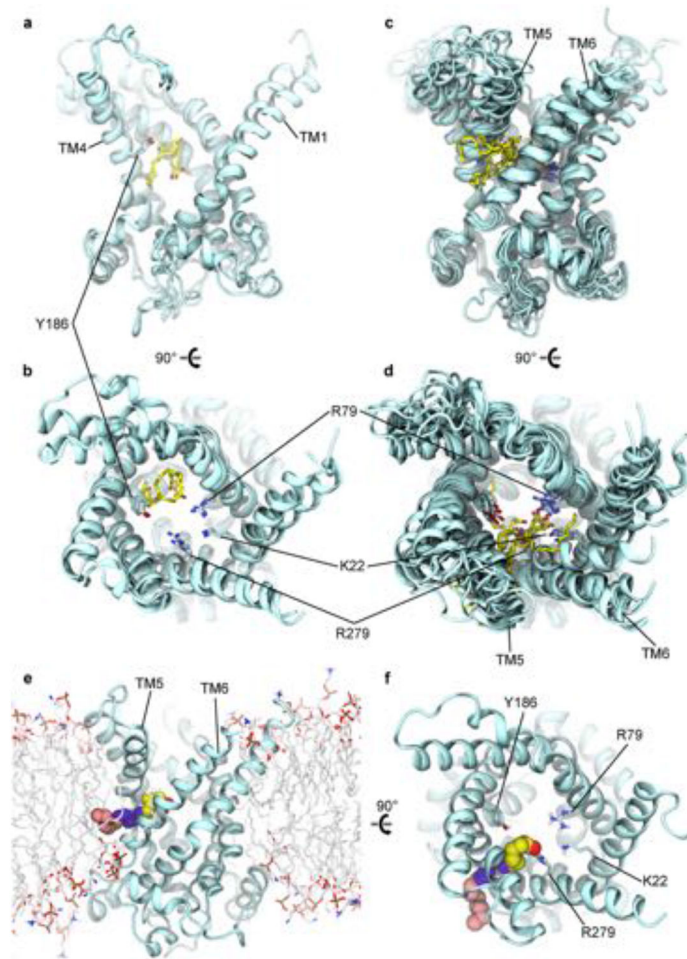
Extended Data Fig. 7 l. Docking reveals a common binding site for small molecules and nucleotide in both c- and m-states of AAC1.

a–j, Final docking poses to the AAC1 c-state (**a–e**) and m-state (**f–j**) structures of deprotonated forms of DNP, FCCP, BAM15, and SF6847, as well as ADP, respectively. In **e** and **j** ADP phosphate groups are hidden for clarity but were included in the docking. The same protein sidechains are shown as sticks in all panels.

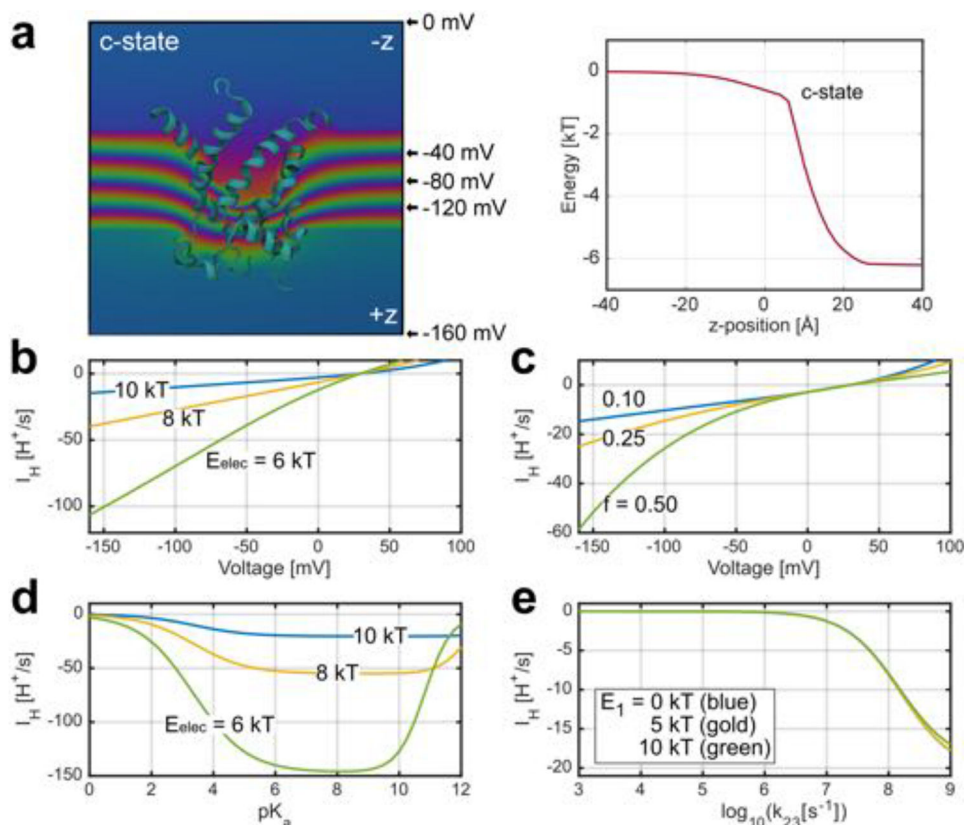


Extended Data Fig. 8 l. DNP binds to the c-state of AAC1 in MD simulations.

a, illustration of the simulation setup used to assess binding of negatively charged DNP to the c-state of AAC1. DNP is shown in sphere representation in the aqueous region of the simulation box. AAC1 is shown as a cyan ribbon, with TM1 and TM6 hidden, and lipid molecules are shown as sticks. The C ζ atom of AAC1 residue R234 at the base of the c-state cavity is shown as a black sphere. Binding was tracked in **b–d** by monitoring the distance from the DNP centre of mass to AAC1 R234 C ζ atom indicated by the dotted line. **b**, Trajectories of negatively charged DNP initially placed in solution far from the binding site. **c**, Trajectories of neutral DNP, initially placed in the binding site. **d**, Trajectories of negatively charged DNP initially in the binding site, with an applied -160 mV membrane potential. In each plot, the two different colored traces are measurements made from two independent simulation trajectories. Simulations in **c** were initiated from docking poses, while those under an applied -160 mV potential in **d** were initiated from the final snapshots of the DNP binding simulations in **b**.



Extended Data Fig. 9 I. Fatty acids bind to the c-state of AAC1 via a TM5/6 fenestration.
a, Arachidonic acid transiently bound to the AAC1 region identified as the DNP/small molecule binding site in contact with protein residue Y186; snapshots are from two independent simulation trajectories. View is from the membrane with TM5 & TM6 of AAC1 hidden to show the cavity. **b**, Top view of structures in panel **a** viewed from the cytoplasm. **c**, Final states of four arachidonic and three palmitic acids bound to the fenestration between AAC1 helices TM5 & TM6; snapshots are from 7 independent simulation trajectories. **d**, Top view of structures in panel **c**. In panels **a–d**, AAC1 is shown as a cyan ribbon and FAs are shown as sticks with yellow carbon and red oxygen atoms. **e**, Side view and **f**, top view of a single structure from panels **c** and **d** with FA atoms shown as spheres and bilayer lipids shown as sticks. Carbon atoms 1–6, 7–12, and 13–20 of the arachidonic acid are colored yellow, blue, and pink, to highlight the parts that are inside the AAC1 cavity, in the TM5/TM6 fenestration, and interacting with bilayer lipids, respectively.



Extended Data Fig. 10 l. Calculation of membrane potential profiles and mathematical model results for AAC1.

a. Membrane potential profile ($V(z)$) across c-state (left) and energy values along the central pore for a monovalent cation ($|e| \cdot V(z)$) (right). The profiles were computed under a -160 mV applied voltage using Poisson-Boltzmann theory (see Materials and Methods) on a configuration extracted from the equilibrated MD simulations. Colour gradients represent isocontours of linearly spaced scalar values ranging from 0 mV (upper region of graphic) to -160 mV (lower region). The water filled cavity facing upward supports a minor fraction of the field with the majority of the field focused across the closed matrix gate of the c-state. These values are quantified in the right panel showing the energy of interaction for a monovalent cation moving through the membrane field. **b–e.** Steady state single-transporter currents as a function of voltage (**b, c**), pK_a (**d**), and rate k_{23} of proton transfer from the uncoupler to matrix pathway (**e**) for the 3-state model shown in Fig. 4g using values in Supplementary Table 2 unless otherwise indicated in the panel. The electrostatic potential in the cavity (E_{elec}) was varied: 10 $k_B T$ (blue), 8 $k_B T$ (gold), 6 $k_B T$ (green) (**b** and **d**). The fraction of the membrane voltage traversed by a H^+ entering from the cytosol was varied: $f = 0.1$ (blue), 0.25 (gold), and 0.5 (green) (**c**). The energy of the proton in the matrix pathway (E_1) was varied: 0 $k_B T$ (blue), 5 $k_B T$ (gold), 10 $k_B T$ (green) (**e**).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgements

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Data availability

All data and materials are available on request from the corresponding authors.

Code availability

The MATLAB code for the mathematical model, force field and docking parameters for DNP, FCCP, BAM15, SF6847, CATR and the homology model of m-state bovine AAC1 are available at Zenodo (<https://doi.org/10.5281/zenodo.5058463>).

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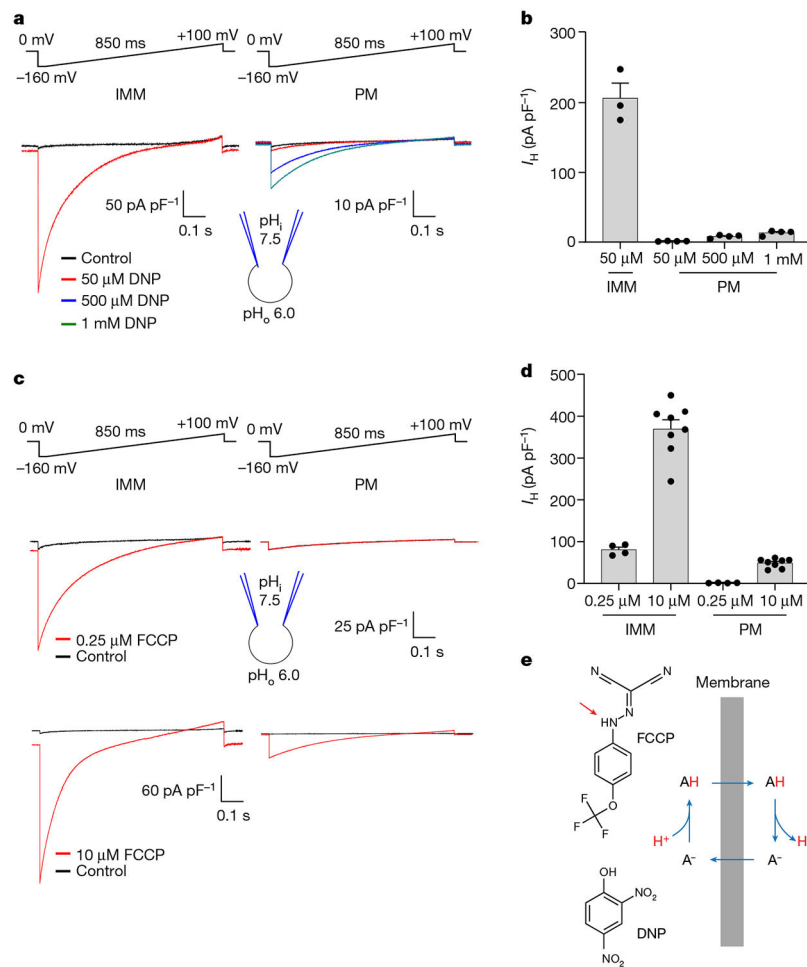


Fig. 1 | H⁺ currents activated by DNP and FCCP in the IMM versus plasma membrane.
a, DNP-induced I_H recorded across the whole IMM of skeletal muscle (IMM, left) or the whole plasma membrane of HEK293 cells (PM, right). **b**, Densities of DNP-induced I_H in the IMM ($n = 3$) versus PM ($n = 4$ for each concentration), measured at -160 mV. Data are mean \pm s.e.m. **c**, FCCP-induced I_H recorded across the whole IMM of skeletal muscle (left) or the whole plasma membrane of HEK293 cells (right). **d**, Densities of FCCP-induced I_H in the IMM ($n = 4$ and $n = 8$) versus PM ($n = 4$ and $n = 8$), measured at -160 mV. Data are mean \pm s.e.m. **e**, Protonatable groups of DNP ($pK_a = 4.1$) and FCCP ($pK_a = 6.2$) indicated by red arrows (left). The mechanism of I_H induced by protonophores (right). ‘A’ indicates protonatable weak acid.

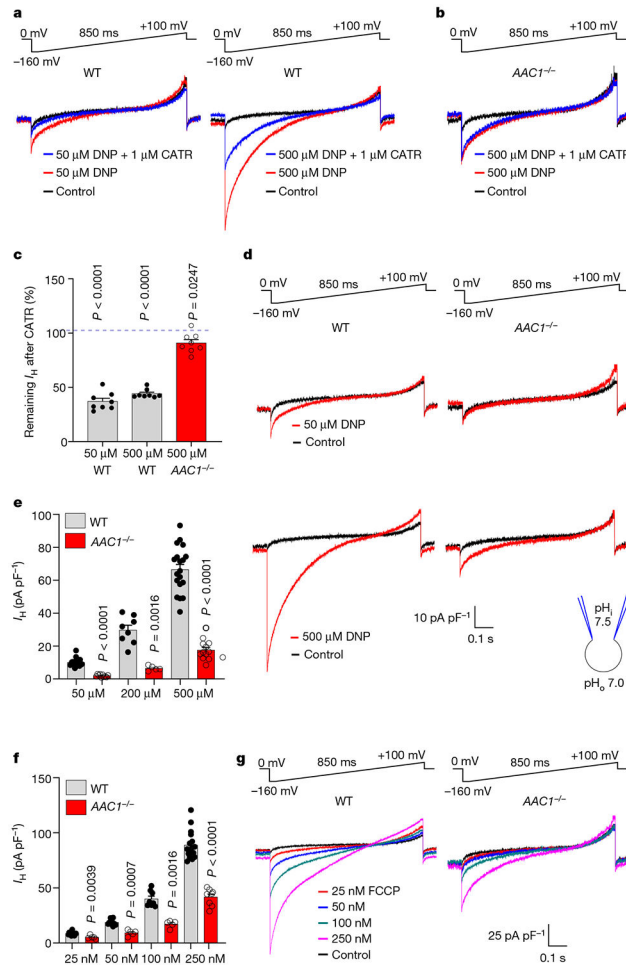


Fig. 2 | DNP and FCCP activate I_H through AAC.

a, I_H induced by 50 μ M (left) or 500 μ M DNP (right) in heart IMM was inhibited by 1 μ M CATR. **b**, I_H induced by 500 μ M DNP in $AAC1^{-/-}$ heart IMM before and after application of 1 μ M CATR. **c**, The remaining fraction of DNP-induced I_H at -160 mV after inhibition by 1 μ M CATR in WT and $AAC1^{-/-}$. Data are mean \pm s.e.m. Statistical analysis was performed using two-tailed paired t -test, compared with I_H before CATR treatment. $n = 8$ for all conditions. **d**, I_H in WT and $AAC1^{-/-}$ heart IMM induced by 50 μ M and 500 μ M DNP. **e**, I_H densities at -160 mV in WT and $AAC1^{-/-}$ heart IMM at different DNP concentrations. Data are mean \pm s.e.m. Statistical analysis was performed using two-tailed Mann-Whitney U -tests, WT versus $AAC1^{-/-}$. $n = 21$ (WT, 50 μ M) and $n = 10$ ($AAC1^{-/-}$, 50 μ M); $n = 8$ (WT, 200 μ M) and $n = 5$ ($AAC1^{-/-}$, 200 μ M); and $n = 19$ (WT, 500 μ M) and $n = 13$ ($AAC1^{-/-}$, 500 μ M). **f**, I_H densities at -160 mV in WT and $AAC1^{-/-}$ heart IMM at different FCCP concentrations. Data are mean \pm s.e.m. Statistical analysis was performed using two-tailed Mann-Whitney U -tests, WT versus $AAC1^{-/-}$. $n = 12$ (WT, 25 nM) and $n = 5$ ($AAC1^{-/-}$, 25 nM); $n = 10$ (WT, 50 nM) and $n = 5$ ($AAC1^{-/-}$, 50 nM); $n = 8$ (WT, 100 nM) and $n = 5$ ($AAC1^{-/-}$, 100 nM); and $n = 17$ (WT, 250 nM) and $n = 8$ ($AAC1^{-/-}$, 250 nM). **g**, I_H induced by 25 nM, 50 nM, 100 nM and 250 nM FCCP in WT and $AAC1^{-/-}$ heart IMM.

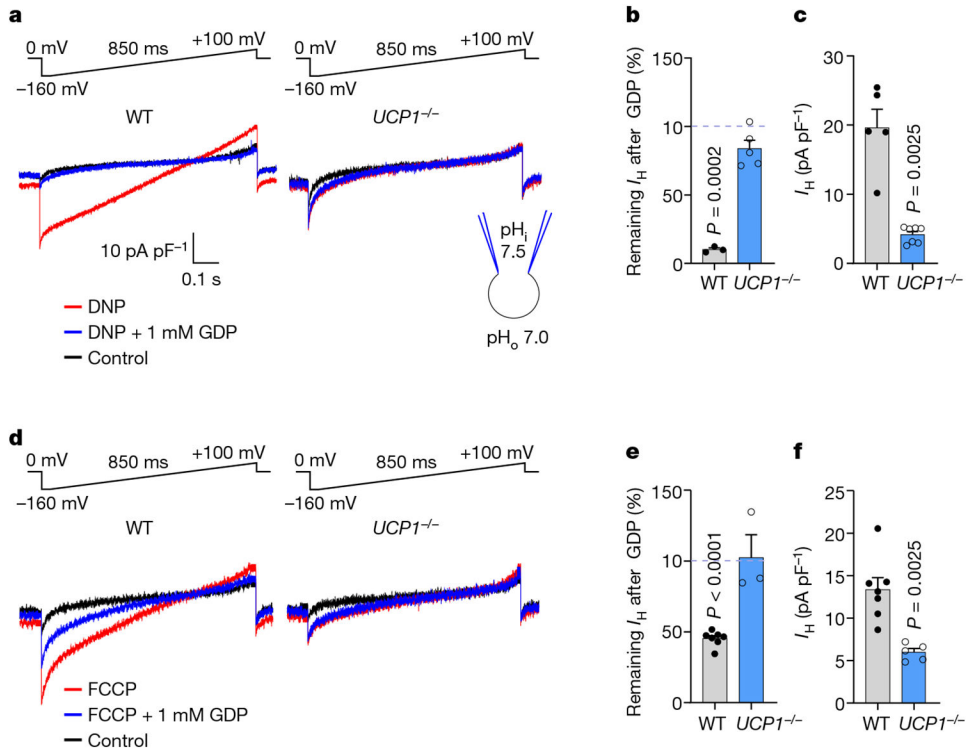


Fig. 3 | DNP and FCCP induce I_H through UCP1.

a, DNP-induced I_H across the IMM of brown fat in WT and *UCP1^{-/-}* before and after application of 1 mM GDP. DNP (500 μ M) was applied in the presence of 10 mM M β CD to extract endogenous membrane FAs. **b**, The remaining fraction of DNP-induced I_H at -160 mV after inhibition by 1 mM GDP in WT ($n = 3$) and *UCP1^{-/-}* ($n = 5$). Statistical analysis was performed using two-tailed paired *t*-tests, I_H without GDP versus with GDP in the WT. **c**, The densities of DNP-induced I_H in WT ($n = 5$) and *UCP1^{-/-}* ($n = 7$) as measured at -160 mV. Data are mean \pm s.e.m. Statistical analysis was performed using two-tailed Mann-Whitney *U*-tests, WT versus *AAC1^{-/-}*. **d**, FCCP-induced I_H across the IMM of brown fat in WT and *UCP1^{-/-}* before and after application of 1 mM GDP. FCCP (10 μ M) was applied in the presence of 10 mM M β CD to extract endogenous membrane FAs. **e**, The remaining fraction of FCCP-induced I_H at -160 mV after inhibition by 1 mM GDP in WT ($n = 7$) and *UCP1^{-/-}* ($n = 3$). Statistical analysis was performed using paired two-tailed *t*-tests, I_H without GDP versus with GDP in the WT. **f**, The densities of DNP-induced I_H in WT ($n = 7$) and *UCP1^{-/-}* ($n = 5$) mitoplasts. I_H was measured at -160 mV. Data are mean \pm s.e.m. Statistical analysis was performed using two-tailed Mann-Whitney *U*-tests, WT versus *AAC1^{-/-}*.

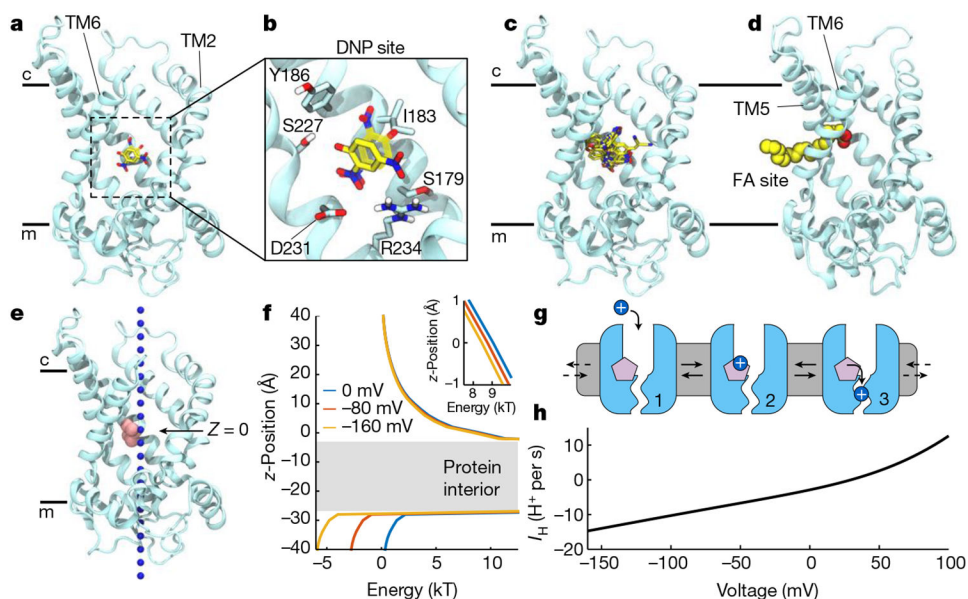


Fig. 4 |. Uncouplers and FAs bind within AAC.

a. Binding poses of deprotonated DNP in the cavity of the AAC1 c-state obtained from docking (solid) and a 500 ns molecular dynamics simulation (transparent) viewed from the membrane with TM1 hidden. AAC1 (cyan) is represented as ribbons, and DNP as sticks (carbon (yellow), oxygen (red) and nitrogen (blue)). **b.** Expanded view of DNP poses in **a.** **c.** Superposition of predicted binding poses for protonated and deprotonated forms of DNP, FCCP, BAM15 and SF6847. **d.** Arachidonic acid from a molecular dynamics simulation showing the carboxylate headgroup occupying the central cavity and aliphatic tail extending into the bilayer between TM5 and TM6. **e.** Hypothetical H_3O^+ (blue spheres) pathway from the intermembrane space (c) to the matrix (m). Pink spheres are titratable groups of docked uncouplers shown in c set to $z = 0$. **f.** Electrostatic energy profile computed along pathway in e using Poisson–Boltzmann continuum electrostatics. The grey region indicates high electrostatic barrier due to lack of solvation in the protein interior. The profiles correspond to applied voltages of 0 mV (blue), -80 mV (red) and -160 mV (yellow). Inset: magnified view of the profile near protonatable groups at $z = 0$. **g.** Cartoon model of the three-state H^+ transport cycle: (1) the uncoupler (pink) bound to c-state of AAC1 (light blue); (2) cytoplasmic H^+ (blue sphere) binds to the uncoupler; and (3) H^+ transitions to the matrix pathway. H^+ exits AAC1 to the matrix as the cycle resets back to state 1. Each step is reversible, but only H^+ movement from the cytosol to matrix is shown because it is favoured under negative matrix voltages. **h.** The steady state single-transporter current–voltage relationship predicted from the model in **g** using the values in Supplementary Table 2 for a DNP-like uncoupler with cytosolic pH = 7 and matrix pH = 7.5. Model equations are provided in Supplementary Note 1.