Inhibition of the hERG potassium channel by phenanthrene: a polycyclic aromatic hydrocarbon pollutant

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Abstract
The lipophilic polycyclic aromatic hydrocarbon (PAH) phenanthrene is relatively abundant in polluted air and water and can access and accumulate in human tissue. Phenanthrene has been reported to interact with cardiac ion channels in several fish species. This study was undertaken to investigate the ability of phenanthrene to interact with hERG (human Ether-a-go-go-Related Gene) encoded Kv11.1 K⁺ channels, which play a central role in human ventricular repolarization. Pharmacological inhibition of hERG can be proarrhythmic. Whole-cell patch clamp recordings of hERG current (IhERG) were made from HEK293 cells expressing wild-type (WT) and mutant hERG channels. WT IhERG was inhibited by phenanthrene with an IC₅₀ of 17.6 ± 1.7 µM, whilst IhERG₁a/₁b exhibited an IC₅₀ of 1.8 ± 0.3 µM. WT IhERG block showed marked voltage and time dependence, indicative of dependence of inhibition on channel gating. The inhibitory effect of phenanthrene was markedly impaired by the attenuated inactivation N588K mutation. Remarkably, mutations of S6 domain aromatic amino acids (Y652, F656) in the canonical drug binding site did not impair the inhibitory action of phenanthrene; the Y652A mutation augmented IhERG block. In contrast, the F557L (S5) and M651A (S6) mutations impaired the ability of phenanthrene to inhibit IhERG, as did the S624A mutation below the selectivity filter region. Computational docking using a cryo-EM derived hERG structure supported the mutagenesis data. Thus, phenanthrene acts as an inhibitor of the hERG K⁺ channel by directly interacting with the channel, binding to a distinct site in the channel pore domain.

Keywords Hydrocarbon · KCNH2 · PAH · Phenanthrene · Pollutant · Potassium channel

Introduction

Humanity’s dependence on fossil fuels has led to the near-ubiquitous pollution of the environment with petrochemicals. Exposure to polluted air is associated with a range of adverse cardiovascular events including heart attacks, strokes and irregular heart rhythms, particularly in people already at risk for these conditions [1]. The mechanisms of cardiotoxicity are complex and can be attributed in part, to nanoparticles containing polycyclic aromatic hydrocarbons (PAHs) that are produced during combustion of coal, petroleum, diesel, tobacco and by cooking food [1]. The adverse health effects of PAHs have long been recognized, but with a focus primarily on carcinogenic compounds, such as benzo(a)pyrene (BAP) [2] and other large molecular weight members (i.e. 4+ rings) of the PAH family. In contrast, the noncarcinogenic smaller PAHs have largely been ignored by the human biomedicine community. These smaller 2- and 3-ring compounds are several orders of magnitude more abundant than the larger 4-ring + PAHs in the fossil fuel sources that feed urban air pollution [3], but are not routinely measured in air pollution; when they are, their ubiquity and abundance are clear. Other than 2-ringed naphthalenes, phenanthrene with 3-rings and its alkylated homologs make up the largest fraction of the measured PAHs in virtually all fuel-related sources. This includes fine particulate matter (e.g. PM₂.₅), such as diesel exhaust particles and broadly collected urban PM₂.₅ (e.g. [4, 5]). More importantly, phenanthrenes are...
volatile so are even more abundant in the vapor phase of vehicle exhaust and urban air (e.g. [6, 7]).

Although the phenanthrenes (and other smaller PAHs) have not been considered fully in the etiology of air pollution-related cardiovascular disease in humans, for nearly three decades they have been a major focus of the cardiotoxicity syndrome that occurs in developing fish following aquatic oil spills (reviewed in Ref. [8]). Oil spills, such as the 1989 Exxon Valdez tanker grounding and the 2010 Deepwater Horizon wellhead blowout released large quantities of PAHs directly into the marine environment, bringing global attention to the cardiotoxic effects of PAHs for fish [1, 9, 10]. Exposure of zebrafish embryos to 3-ringed PAHs was found to produce bradycardia and arrhythmia consistent with atrioventricular block; the authors suggested Ether-à-go-go-Related Gene (ERG) encoded K\(^+\) channels as a potential target that might mediate such effects [9]. Crude oil extracts from the Deepwater Horizon disaster were then shown to exert direct effects on the function of tuna cardiac myocytes: they produced impairment of calcium cycling and action potential prolongation, associated with inhibition of the rapid delayed rectifier K\(^+\) current (I\(_{Kr}\), the ion current carried by erg [11]). Subsequent work isolated the 3-ringed PAH phenanthrene as the key moiety involved in these effects producing a concentration-dependent ventricular action potential prolongation and inhibition of I\(_{Kr}\) [12]. Phenanthrene has subsequently been shown to increase sodium current and reduce both calcium current and I\(_{Kr}\) in ventricular myocytes from rainbow trout [13], with ventricular action potential prolongation and reductions in calcium current and I\(_{Kr}\) with phenanthrene also recently reported for brown trout cardiomyocytes [14]. Furthermore, phenanthrene has recently been reported to inhibit calcium current and I\(_{Kr}\) in zebrafish cardiomyocytes [15]. There is debate as to whether toxic effects of pollutants on fish from crude oil may largely result from membrane disruption rather than interaction with specific receptors (e.g. [16]). Whilst the effects of phenanthrene on specific cardiac ion channel currents may be consistent with a direct rather than nonselective effect, definitive evidence for this is currently lacking.

The reported effects of phenanthrene on fish I\(_{Kr}\) are of relevance to human cardiovascular health because I\(_{Kr}\) is known to be of key importance not only in animal model species, but also to normal human ventricular repolarization [17, 18]. I\(_{Kr}\) channels are comprised of tetramers of pore-forming Kv11.1 subunits encoded by hERG (human Ether-à-go-go-Related Gene, alternative nomenclature KCNH2; [19, 20]). Loss- and gain-of function mutations to hERG, respectively underlie forms of inherited long and short QT syndromes (LQTS and SQTS, respectively) [19, 21]. Moreover, due to unique structural features, hERG channels are highly susceptible to pharmacological blockade, leading to a drug-induced form of LQTS, with its associated risk of dangerous ventricular arrhythmias [17, 18]. Phenanthrene is relatively abundant in polluted air and water. It is lipophilic and can access and accumulate in human tissue via skin and mucus membranes [1]. Given the known susceptibility of hERG to pharmacological blockade and the fact that fish I\(_{Kr}\) is inhibited by phenanthrene, it is reasonable to expect that hERG itself may be susceptible to phenanthrene inhibition. Furthermore, alignment of fish ERG and hERG sequences reveals a high level of identity in regions of the channel that constitute the canonical drug binding site (Fig. 1A and [1]). It therefore seems likely that human I\(_{Kr}/hERG\) also may be susceptible to inhibition by phenanthrene. Accordingly, this study was undertaken: (i) to determine the propensity or otherwise of phenanthrene to inhibit ionic current (I\(_{hERG}\)) carried by recombinant hERG channels; (ii) to explore the underlying mechanism of any observed effect. Some of this work has been published in meeting abstract form [22, 23].

Materials and methods

Maintenance of mammalian cell lines and cell transfection

As in previous studies (e.g. [24–26]), experiments were performed on HEK293 cells stably expressing WT hERG [27] or transiently transfected with hERG mutant cDNAs. The hERG stable line was generously provided by Professor Craig January [27]. HEK293 cells (ECACC, Porton Down, UK) were transiently transfected with cDNA plasmids using Lipofectamine 2000 (Invitrogen, Paisley, UK) according to the manufacturer’s instructions. Expression plasmid encoding CD8 was also added (in pIRES, donated by Dr I Baró, University of Nantes, France) as a marker for successful transfection. Cells were passaged using enzyme-free cell dissociation solution (Millipore, Watford, UK) and plated onto sterilized shards of glass coverslips in 40-mm petri dishes containing a modification of Dulbecco minimum essential medium with Glutamax-1 (DMEM; Invitrogen, Paisley, UK). This was supplemented with 10% fetal bovine serum (Gibco, Gloucester, UK), 50 μg/mL gentamycin (Invitrogen, Paisley, UK) and 400 μg/mL geneticin (G418, Invitrogen, Paisley, UK) for the WT hERG-expressing line. For experiments utilizing cells transiently transfected with hERG mutants, recordings were performed 24–72 h after transfection. Successfully transfected cells (positive to CD8) were identified using Dynabeads \textsuperscript{®} (Invitrogen, Paisley, UK) [25, 26]. The N588K, S624A, Y652A, M651A and F557L mutations to hERG1a have all been used in prior studies from our laboratory (e.g. [24–26, 28, 29]), as has hERG1a/1b co-expression [25, 30]. Zebrafish ERG (zERG) was synthesised and provided in pcDNA3.1 by Genscript (Leiden, The Netherlands; NCBI Reference Sequence: NM_212837.1).
The F656V mutation was made to hERG1a as described in Ref. [31]. F656T was made using QuikChange (Agilent) mutagenesis using conditions described in Ref. [31] and the following primer sequences:

forward-5'-GTATGCTAGCATCACCGGCAACGT
reverse-5'-CGACACGTTGCGGGTGATGCTAGC

ATAC-3'
Experiments were performed on hERG1a current (\( I_{\text{hERG1a}} \)) except for the data in Fig. 2B, C, which were conducted using co-expressed hERG1a and 1b channels (\( I_{\text{hERG1a/1b}} \)) and Supplementary Fig. 2 which shows data from zERG (\( I_{\text{zERG}} \)).

**Electrophysiological recording**

Recordings were made as described previously [25, 26]. In brief, electrophysiological recordings were made using an Axopatch 200B amplifier (Molecular Devices) with a CV-4/100 headstage and data acquisition via a Digidata 1320 interface (Molecular Devices). Glass shards with plated HEK293 cells were placed in the recording chamber of an inverted microscope (Nikon Diaphot, USA). The extracellular superfusate was a standard Tyrode’s solution containing (in mM): 140 NaCl, 4 KCl, 2.0 CaCl2, 1 MgCl2, 10 glucose and 5 HEPES (titrated to pH 7.4 with NaOH) [24–26]. Patch pipettes (AM systems Inc, USA) had resistances of 2–4 MΩ and were filled with a solution containing (in mM): 130 KCl, 1 MgCl2, 5 EGTA, 5 MgATP and 10 HEPES (titrated to pH 7.2 with KOH) [24–26]. Series resistance was typically compensated by 60–80%. Currents were filtered at 1–5 kHz depending on the voltage protocol used and were digitized at 10 kHz. Except for online

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**Fig. 2 A** Bar charts comparing deactivation parameters for \( I_{\text{hERG}} \) (hERG1a) tails elicited on repolarization to −40 mV from +20 mV in Control (black) and 10 µM phenanthrene (grey). Biexponential fitting yielded fast and slow deactivation time constants (\( \tau_f \) and \( \tau_s \), respectively), plotted in 

**Bi** shows representative traces of hERG1a/1b \( I_{\text{hERG}} \) elicited by the standard protocol shown in Fig. 1 in the absence and presence of 10 µM phenanthrene. 

**Bi** shows concentration response data for 5 phenanthrene concentrations (1 µM, 5 µM, 10 µM, 30 µM; 100 µM; \( n \)= at least 5 at each concentration). The data were fitted with Eq. 2, yielding a half-maximal inhibitory concentration (IC\(_{50}\)) of \( 1.8 \pm 0.3 \) µM and Hill coefficient of \( 0.80 \pm 0.09 \). 

**Ci** shows that the proportion of fast deactivating current increased in the presence of phenanthrene (* \( p < 0.05 \) \( n = 6 \)). Bar charts show mean ± SEM values, with individual experimental values in control and phenanthrene superimposed as filled circles and squares respectively.
Supplementary Fig. 3A, all measurements were made at room temperature (21 ± 1 °C) and run-down correction was not performed. Room temperature was employed to facilitate accurate evaluation of phenanthrene effects because in pilot experiments at 37 °C, stability of phenanthrene-containing superfusate appeared variable. It also facilitated comparison of phenanthrene blocking potency between human and zebrafish ERG data in this study and with prior recordings from fish myocytes conducted at room temperature [11–13].

**Trafficking assay**

Evaluation of the effects of 3 and 30 µM phenanthrene on hERG channel trafficking was performed using a LI-COR based In-Cell/On-Cell Western assay, as described previously [32] (see online supplement for more details).

**Phenanthrene**

Phenanthrene was obtained from Merck (Sigma-Aldrich) and dissolved in DMSO to give stock solutions of 20, 30 and 50 mM. Experimental solutions with phenanthrene concentrations shown in the Results contained a maximum of 0.1% DMSO.

**Computational docking**

Docking of phenanthrene to hERG was initially performed using the cryo-EM derived structure for hERG [33] (PDB code 5VA2), as described previously [26, 29, 34]. A model closely related to the cryo-EM structure was obtained from a short molecular dynamics (MD) simulation in which the F656 side chain of one of the four hERG subunits was found to reorient towards the pore—this subunit was then replicated around all four pore subunits to produce a model with all four F656 side chains facing the pore; this structure was used for the docking data shown in Fig. 6. The phenanthrene structure was converted from SMILE representation (obtained from PubChem database) to a 3D structure, hydrogens were added and the molecule energy minimised.

Phenanthrene was docked in the hERG structure and in the related model using GOLD (GOLD version 5.6; Cambridge Crystallographic Data Centre, Cambridge, UK). The central pore cavity was initially chosen as a binding site where a radius of 10 angstrom extended from the centre of the cavity and in a level with a middle point between the canonical aromatic residues F656 and Y652. The side chains of these aromatic residues were set to be freely flexible during docking simulations. Rotamer sampling was maximally set to 300,000 generations. Dockings were scored by Goldscore and rescored by Chemscore. Two hundred docking repeats were made in each case and the low-energy-score poses were selected and inspected. Phenanthrene was also docked within a side pocket under the selectivity filter in the open pore F656-rotated hERG model. This binding pocket was centred above the β-carbon of Y652 and encompassed a volume having a radius of 7 angstrom.

Within this selection, the side chains for the following residues from subunit A of the tetrameric channel were permitted to rotate freely: F557, L622, T623, S624, L650, M651, Y652, I655. F656 side chains from adjacent subunits (A and B) of the channel were also allowed to rotate freely. Similar setting and parameters were used as above where also 200 docking repeats for each drug were generated and low energy poses identified. In all docking runs the maximal side chain rotamer sampling allowed in GOLD was used (free rotation of 10 selected residue side chain as well as sampling of Ser, Thr and Tyr side chain hydroxyl rotamers to optimize hydrogen bond interactions) with the specific set of flexible residues chosen to match the specific binding location tested.

The results were presented using PyMOL Molecular Graphics System, Version 2.0 Schrödinger, LLC.

**Data presentation and analysis**

The data are presented as mean ± SEM of the number of independent experiments indicated (n). Statistical comparisons were made using a Student’s t test, one- or two-way analysis of variance (ANOVA) followed by a Bonferroni or Dunnett’s post test, as appropriate. *P* values <0.05 were considered to be statistically significant.

Fractional block of hERG current (I_{hERG}) by the different phenanthrene (Phen) concentrations studied was determined using the equation:

\[
\text{Fractional block} = 1 - \left( \frac{I_{\text{hERG-Phen}}}{I_{\text{hERG-Control}}} \right)
\]  \hspace{1cm} (1)

where “Fractional block” refers to the degree of inhibition of hERG current by a given concentration of phenanthrene. I_{hERG-Phen} and I_{hERG-Control} represent current amplitudes in the presence and absence of phenanthrene.

Concentration–response data were fitted by a standard Hill equation of the form:

\[
\text{Fractional block} = 1 - \left( \frac{1}{1 + \left( \frac{\text{IC}_{50}}{[\text{Phen}]} \right)^h} \right)
\]  \hspace{1cm} (2)

where IC_{50} is [Phen] producing half-maximal inhibition of the I_{hERG} tail and *h* is the Hill coefficient for the fit.

Half maximal activation voltages for I_{hERG} were obtained from current–voltage (I–V) relations from I_{hERG} tails measured at −40 mV in the absence or presence of phenanthrene following voltage commands to different test potentials, using the following Boltzmann equation:

\[
I = I_{\text{max}} \left( 1 + \exp \left( \frac{V - V_{0.5}}{k} \right) \right)
\]  \hspace{1cm} (3)

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where I = \( I_{\text{hERG}} \) tail amplitude following test potential \( V_m \), \( I_{\text{max}} \) is the maximal \( I_{\text{hERG}} \) tail observed during the protocol, \( V_{0.5} \) is the half maximal activation voltage of \( I_{\text{hERG}} \) and \( k \) is the slope factor describing \( I_{\text{hERG}} \) activation.

Voltage-dependent activation curves were constructed by calculating activation variables at 2 mV intervals between −80 mV and +40 mV. Values for \( V_{0.5} \) and \( k \) were derived from experimental fits to I–V data using Eq. (3) were inserted into the following equation:

 Activation parameter \( = \frac{1}{1 + \exp \left( \frac{(V_{0.5} - V_m)}{k} \right)} \)  

where the ‘activation parameter’ at test potential \( V_m \) lies between 0 and 1 and \( V_{0.5} \) and \( k \) have the meanings described above for Eq. 3.

### Results

**WT \( I_{\text{hERG}} \) inhibition by phenanthrene**

The sensitivity of \( I_{\text{hERG}} \) to inhibition by phenanthrene was determined using the protocol shown in Fig. 1B. This was comprised of a 2 s depolarization from −80 mV to +20 mV, followed by repolarization to −40 mV, at which the resurgent tail current that is typical of hERG was observed. \( I_{\text{hERG}} \) tail magnitude was measured relative to current elicited by a brief (50 ms) pulse from −80 to −40 mV that preceded the 2 s voltage command. This protocol has been used in multiple prior studies of \( I_{\text{hERG}} \) pharmacology from our laboratory (e.g. [24–26, 35]). Figure 1B shows exemplar records of \( I_{\text{hERG}} \) in control solution, the presence of 10 μM phenanthrene and following washout, whilst Fig. 1C shows a continuous plot of \( I_{\text{hERG}} \) tail amplitude during the experiment from which these example traces were taken. Phenanthrene application led to a progressive decline in \( I_{\text{hERG}} \) amplitude, which was largely reversible on washout. Six concentrations of phenanthrene, ranging from 1 to 100 μM, were tested in similar experiments. Higher concentrations could not be investigated due to issues with phenanthrene solubility and exposure of cells to higher levels of solvent. Fractional inhibition of \( I_{\text{hERG}} \) tails at each phenanthrene concentration was ascertained using Eq. 1 and values from different experiments at each concentration pooled to produce the concentration response plot shown in Fig. 1D. These data were fitted with Eq. 2, which yielded a half-maximal inhibitory concentration (IC\(_{50}\)) of 17.6 ± 1.7 μM and Hill slope of 0.94 ± 0.09. At the highest concentration tested (100 μM), the mean level of tail current inhibition attained was 79.1 ± 2.9% (\( n = 5 \)). It is apparent from the traces shown in Fig. 1B that in addition to reducing \( I_{\text{hERG}} \) amplitude, phenanthrene altered the deactivation time course of the \( I_{\text{hERG}} \) tail. This effect was quantified by standard biexponential fitting of deactivating \( I_{\text{hERG}} \) tails at −40 mV following 2 s duration commands to +20 mV. As shown in Fig. 2A, both fast and slow time constants of deactivation (\( \tau_f \) and \( \tau_s \)) were significantly decreased in the presence of phenanthrene, whilst the proportion of deactivation described by \( \tau_f \) increased. These findings indicate that phenanthrene application resulted in significant acceleration of \( I_{\text{hERG}} \) deactivation.

### Inhibition by phenanthrene of \( I_{\text{hERG}} \) elicited at different voltages

Voltage dependence of hERG inhibition was investigated for hERG1a using the I–V protocol shown in Fig. 3A; although there was a trend towards increased \( I_{\text{hERG}} \) block at the higher temperature this did not attain statistical significance (\( p > 0.05 \)).
This protocol was similar to that used in Fig. 1, but incorporated progressively larger depolarizations (in 10 mV increments) to test potentials between −30 and +60 mV. Whilst both pulse and tail \( I_{\text{hERG}} \) with voltage commands to more positive membrane potentials in the range tested were reduced by phenanthrene (10 µM), at more negative voltages in the tested range, the current appeared to increase in amplitude in the presence of phenanthrene. Figure 3B, C shows normalized I–V relations for end-pulse current (Fig. 3B) and tail current (Fig. 3C). As Fig. 3B shows the end pulse I–V relation showed the bell-shaped characteristic of \( I_{\text{hERG}} \) in both control and phenanthrene containing solutions. Superimposed on the same voltage axes is a plot of fractional block of \( I_{\text{hERG}} \) at each voltage. As is clearly illustrated, at potentials below +10 mV facilitation (negative plotted values, representing an increase in current) of \( I_{\text{hERG}} \) was observed. This gave way to a progressive increase in end pulse current block, up to ~ +40 mV. Normalized \( I_{\text{hERG}} \) tails were plotted and fitted with Eq. 4 in Fig. 3C, along with a superimposed plot of fractional inhibition (cf [25, 35, 42]). For all test potentials negative to ~ 0 mV, facilitation (a fractional increase) in \( I_{\text{hERG}} \) tail amplitude at -40 mV was seen, whilst fractional inhibition increased markedly up to ~ +20 mV, then levelled off at positive potentials. The normalized \( I_{\text{hERG}} \) tail I–V relation exhibited a left-ward voltage shift in the presence of phenanthrene. In control solution the \( V_{0.5} \) value obtained was 1.3 ± 1.6 mV \((k = 7.2 ± 0.2 \text{ mV, } n = 16)\), whilst in 10 µM phenanthrene \( V_{0.5} \) was -8.9 ± 1.5 mV \((P < 0.001 \text{ vs Control}; k \text{ unchanged } 7.8 ± 0.3 \text{ mV, } n = 16)\).

Supplementary Fig. 3B shows an alternative display format: plots of \( I_{\text{hERG}} \) tail current density. These illustrate the increase in tail current density at negative voltages and inhibition at more positive voltages in the tested range and also show the leftward shifted activation in phenanthrene. Thus, phenanthrene produced a ~ 10 mV negative shift in voltage-dependent activation associated with the facilitation of \( I_{\text{hERG}} \) at negative voltages. This phenomenon has been observed previously for other hERG-blocking agents, including azimilide [43], 4-aminopyridine [42] and sarizotan [35]. Notably, the steep change in fractional inhibition coincided with the steep portion of the voltage-dependent
activation relation for $I_{\text{hERG}}$ (Fig. 3C) consistent with an activation-dependent inhibitory mechanism.

**Depolarization time and $I_{\text{hERG}}$ inhibition by phenanthrene**

The voltage dependence of $I_{\text{hERG}}$ inhibition by phenanthrene indicates gating dependence of channel inhibition by the compound. To probe this further, we conducted experiments in which the extent of $I_{\text{hERG}}$ inhibition by voltage commands of different durations was established [42, 44, 45]. For these experiments, activating pulses to $+40$ mV (a potential observed to be at the top of the voltage-dependent activation relations for control and phenanthrene) of different durations between 50 and 1100 ms were applied. These were first applied in control superfusate, the cell under study was then rested for 3–5 min whilst being superfused with 30 µM phenanthrene and the protocol was reapplied. The extent of $I_{\text{hERG}}$ inhibition by the compound for the different duration commands was then evaluated by assessing the extent of tail current block. The left panel of Fig. 4A shows a family of current traces in control solution (current—upper traces, voltage protocol—lower traces), whilst the right panel of Fig. 4A shows corresponding data in 30 µM phenanthrene. The increase in tail current amplitude as pulse duration increased (for both conditions) reflects the progressively increased time periods for activation of $I_{\text{hERG}}$. As these example traces indicate, there was little inhibition of $I_{\text{hERG}}$ following brief depolarizations, but inhibition increased with command pulse duration. The mean fractional block data are plotted against command pulse duration in Fig. 4B, showing a clear time dependence of inhibition, with block increasing markedly as the command pulse increased between 50 and 500 ms, levelling out at longer pulse durations in the range tested. Thus, there was clear dependence of the response to phenanthrene on the duration of the activating test command ($p < 0.0001$ one-way ANOVA; $n = 5$). A monoexponential fit to the data in Fig. 4B yielded a time constant for development of inhibition $\tau_{\text{inh}}$ of $130.2 \pm 15.4$ ms. The increase in tail current inhibition as command pulse duration lengthened was associated with an apparent increase in rate of development of $I_{\text{hERG}}$ activation at $+40$ mV, with $\tau_{\text{act}}$ values of $123.3 \pm 75.6$ ms and $75.6 \pm 9.8$ ms, respectively, in control and phenanthrene ($p < 0.01$). The results of this experiment are consistent with a clear dependence of phenanthrene inhibition of $I_{\text{hERG}}$ on channel gating, with little or no inhibition of closed channels evident from the plot in Fig. 4B.

**Probing the mechanism of $I_{\text{hERG}}$ Inhibition with mutagenesis**

In order to investigate gating dependence of inhibition further, experiments were conducted using the N588K attenuated-inactivation mutation. The N588 residue is located in the external S5-Pore linker region of the channel; it has utility for investigating inactivation dependence of hERG inhibition [46, 47] as it perturbs inactivation gating without directly modifying channel structure within regions of the pore typically implicated in drug binding. Figure 5Ai shows the effect of 30 µM phenanthrene on WT $I_{\text{hERG}}$, illustrating marked current inhibition by this concentration of the compound. In contrast, only slight inhibition of N588K $I_{\text{hERG}}$ was observed (Fig. 5Aii, mean data in Fig. 5H). This indicates that inactivation competency was required for optimal

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**Fig. 4** A Upper traces show representative $I_{\text{hERG}}$ records elicited by protocol shown as lower traces. Tail currents were elicited at $-40$ mV following depolarizing pulses to $+40$ mV for differing time periods between 50 and 1100 ms. Left hand panel shows recordings in control solution and right-hand panel contains recordings from the same cell during exposure to 30 µM phenanthrene. B Plot of mean fractional block of $I_{\text{hERG}}$ tail by 30 µM phenanthrene against command pulse duration ($n = 5$). A monoexponential fit to these data (from each experiment) gave an inhibition time constant ($\tau_{\text{inh}}$) of $130.2 \pm 15.4$ ms. A one-way ANOVA showed an overall significance in the time dependence of inhibition ($p < 0.0001$), with (Bonferroni) post hoc testing showing that inhibition following pulses of up to 200 ms in duration was significantly different from all durations of 500 ms and longer ($p < 0.05$ or less for all comparisons).
interaction between phenanthrene and its binding site on the hERG channel. The S624 residue, located near the base of the hERG channel selectivity filter, has been implicated in hERG inhibition by a number of compounds (e.g. [48, 49]). Figure 5B shows effects of 30 µM phenanthrene on S624A hERG. Pulse current appeared significantly augmented and tail current amplitude unchanged by phenanthrene with the S624A mutation (mean data are shown in Fig. 5H). This implicates the S624 residue, directly or indirectly, as being important for phenanthrene binding to the hERG channel. We proceeded to investigate the roles of two aromatic residues, Y652 and F656 in phenanthrene action. For virtually all compounds that have been investigated in detail, interactions with one or both of these residues are obligatory for drug binding and channel inhibition [17, 50]. Remarkably, the inhibitory effect of 30 µM phenanthrene on Y652A hERG (Fig. 5C, mean data in Fig. 5H) was significantly greater than on WT hERG. Supplemental Fig. 4A shows mean concentration–response data for Y652A hERG, which yielded an IC50 of 0.46 ± 0.01 µM, Hill slope: 0.58 ± 0.11 (at the highest phenanthrene concentration tested (30 µM) hERG tails were inhibited by 98.7 ± 2.6%; n = 7). The F656V mutation (Fig. 5D, mean data in Fig. 5H) also did not impair the inhibitory effect of phenanthrene, showing a trend towards increased block, albeit that this was statistically insignificant. We also investigated effects of a second mutation at F656:F656T. A small reduction in inhibition of F656T hERG by 30 µM is shown in the exemplar traces shown in Fig. 5E; however, mean pooled data (Fig. 5H) show that this effect was not statistically significantly different from block of WT hERG. Phenanthrene is therefore distinct from most hERG-blocking drugs thus far studied in not requiring aromatic residues at positions 652 and 656 for inhibition to occur. Residue Y652 has previously been implicated in voltage dependence of hERG inhibition (e.g. [51, 52]). In this respect, it is notable that whilst the Y652A mutation increased rather than decreased phenanthrene inhibition, it largely abolished the voltage dependence of inhibition seen for WT hERG (Supplemental Fig. 4B). Residue F557 on the S5 helix of the hERG channel has been identified as a determinant of hERG inhibition and lies adjacent to Y652 on the S6 helix [29, 50, 53, 54]. No inhibition of F557L hERG was observed with 3 concentrations of phenanthrene (between 10 and 100 µM). Figure 5F shows representative traces of effects of 30 µM phenanthrene on F557L hERG. No inhibition was evident, rather the pulse and peak tail current were slightly larger in the presence of phenanthrene than in control solution (mean data shown in Fig. 5H). This observation implicates residue F557 as of critical importance to phenanthrene inhibition (e.g. [51, 52]). The publication of a cryo-EM structure for hERG has shown the presence of 4 hydrophobic pockets that surround the central cavity of the channel, which may provide interaction site(s) for drugs [33]. Lipophilic drug access to the channel pore involving F557 and S6 residue M651 has recently been suggested for the bradycardic agent ivabradine [54]. Figure 5G
shows effects of the M651A mutation on \(I_{\text{hERG}}\) block by 30 \(\mu\)M phenanthrene. In contrast to WT \(I_{\text{hERG}}\) M651A was little affected by phenanthrene (mean data shown in Fig. 5H), implicating this residue in the compound’s inhibitory action.

**Phenanthrene inhibition explored through computational docking**

Low energy score docking poses consistent with mutagenesis data were not obtained for phenanthrene bound into the structure of hERG obtained from cryo electron microscopy (cryo-EM). Phenanthrene was unable to make multiple interactions with aromatic side chains when docked into the canonical pore blocker site below the selectivity filter within the \(K^+\) permeation path (not shown). This results from the absence of F656 side chain conformations in the cryo-EM structure that project the side chains towards the pore and the rigidity of the phenanthrene tri-aromatic ring structure that limits its ability to conform to the arrangement of Y652 side chains within the pore. The rigidity of the structure also limited identification of low energy score configurations for phenanthrene binding within one of the hydrophobic pockets below the pore helix in the cryo-EM structure. In addition, the projection of the F656 side chains towards the S5 helix in the cryo-EM structure limits the localisation of phenanthrene in a configuration that allows interactions with the side chains of F557 and M651.

Low energy poses were obtained for phenanthrene bound within a side pocket of a hERG model obtained by very short MD simulation in which an F656 side chain rotated away from its configuration in the cryo-EM structure towards the pore. A low energy score pose consistent with the mutagenesis data is shown in Fig. 6. Phenanthrene sits deep within a hydrophobic pocket making aromatic stacking interactions with F557 and hydrophobic interactions with M651. This binding mode is facilitated by the rotation of the F656 side chain out of this binding pocket.

**Discussion**

**Potency of observed \(I_{\text{hERG}}\) inhibition and environmental relevance for human health**

Phenanthrene has previously been reported to inhibit native \(I_{Kr}\) from bluefin tuna myocytes at room temperature by nearly 60% at 5 \(\mu\)M and by over 80% at 25 \(\mu\)M [11]. 30 \(\mu\)M phenanthrene inhibited rainbow trout myocyte \(I_{Kr}\) by 79% [13], whilst an IC\(_{50}\) for inhibition of brown trout myocyte \(I_{Kr}\) of 7.2 \(\mu\)M has been reported [14]. Very recent work has indicated that \(I_{Kr}\) from zebrafish ventricular myocytes is inhibited by phenanthrene with an IC\(_{50}\) of 3.3 \(\mu\)M [15], which is in fair agreement with the value of 8.7 \(\mu\)M obtained in this study for \(I_{\text{hERG}}\). We observed IC\(_{50}\) values for \(I_{\text{hERG}}\) carried by hERG1a channels of 17.6 \(\mu\)M and for co-expressed hERG1a/1b of 1.8 \(\mu\)M, which are broadly comparable to the potency of phenanthrene against fish \(I_{Kr}\). It is notable that for \(I_{\text{hERG}}\) 1a/1b, which is thought to underlie native mammalian \(I_{Kr}\) [37–39], the maximal level of observed \(I_{\text{hERG}}\) tail
inhibition (at 100 µM) was near complete (~97%). For \( I_{\text{hERG}} \) carried by hERG1a alone, the maximal level of observed tail current inhibition (at 100 µM) did not exceed 80%. Although the result for \( I_{\text{hERG}1a/1b} \) (and also that for Y652A on a hERG 1a background) indicates that complete or near complete channel inhibition by phenanthrene is possible, we cannot entirely exclude the possibility that maximal possible attainable inhibition of WT hERG1a \( I_{\text{hERG}} \) may be <100%. However, due to solubility concerns it was not possible to test phenanthrene concentrations higher than 100 µM.

Data on plasma levels of PAHs in humans are sparse, in part because convention focuses on other measures—either PAH concentrations per given volume of air or urinary metabolite concentrations. Plasma concentrations of phenanthrene and/or other PAHs in the nM range have been reported in the literature (e.g. [55, 56]). However, as has been highlighted elsewhere, phenanthrene is highly lipophilic and so is likely to exhibit tissue accumulation [1]. This has been demonstrated for waterborne exposures of developing fish embryos in which phenanthrene tissue levels have reached micromolar levels (e.g. [57]) and in similarly substantial accumulations of PAHs in avian tissues [58]. Road pavers and coke plant workers show urinary phenanthrol levels of ~35–686 µg/L (approximately 0.2–3.5 µM; [59]). Such data are suggestive of quite high concentrations in subpopulations who, through occupational risk, are exposed routinely to high levels of PAHs. Moreover, a recent comparison of serum phenanthrene levels in Greek citizens found increased levels in those with heart failure (231 µg/L; ~1.3 µM) compared to those without heart failure (56.5 µg/L; 314 nM) [60]. Thus, it is likely to be those with high PAH exposure and/or those with pre-existing cardiovascular morbidity who will be most susceptible to adverse reactions to phenanthrene.

**Evidence for a direct channel interaction**

Phenanthrene has a simple structure of three fused benzene rings and given its high lipophilicity [1], a natural question is whether it exerts direct interactions with the hERG channel or whether its inhibitory effect is secondary to accumulation of phenanthrene in membrane lipid? Multiple lines of evidence provided here support a direct interaction with the channel. First, the lack of significant effect of phenanthrene on hERG trafficking at 3 and 30 µM (Supplementary Fig. 1), indicates that altered trafficking is unlikely to contribute an additional inhibitory effect on the channel’s function during chronic exposure. Second, the structurally related antimalarial drug halofantrine (3-[(dibutylamino)-1-[1,3-dichloro-6- (trifluoromethyl)phenanthren-9-yl]propan-1-ol), which contains a substituted phenanthrene, is known to inhibit \( I_{\text{hERG}} \) in an open state-dependent fashion, with sensitivity to mutation of S6 helical aromatic residues [61, 62]. Third, in this study phenanthrene inhibition of WT \( I_{\text{hERG}} \) exhibited voltage- and time dependence that demonstrate contingency on channel gating for the compound to access its inhibitory binding site. Fourth, the profound reduction in the \( I_{\text{hERG}} \) inhibitory effect of phenanthrene on inactivation-impaired N588K channels suggests a requirement for intact inactivation gating for optimal access to the inhibitory binding site. Although \( I_{\text{hERG}} \) carried by hERG1a/1b (which showed enhanced sensitivity to phenanthrene inhibition) has been reported to exhibit a modest (+28 mV) positive shift in voltage dependence of inactivation compared to WT hERG alone [39], the inactivation shift produced by the N588K mutation is much greater (+60 to +90 mV; [28, 63]) and therefore more impactful on \( I_{\text{hERG}} \) elicited by a command protocol to +20 mV, as used here to evaluate \( I_{\text{hERG}} \) inhibition. Our data suggest that the profound disturbance of inactivation gating with this mutant adversely affects channel conformation(s) favouring phenanthrene access/binding. Finally, the results of our mutagenesis experiments and docking simulations implicate specific mutations as phenanthrene binding determinants. Thus, the results of this study support a conclusion that phenanthrene acts as a direct hERG channel inhibitor. To our knowledge, the data from this study constitute the most detailed information on the basis of phenanthrene inhibition of any ion channel or electrogenic transporter.

**Facilitation accompanying phenanthrene block of \( I_{\text{hERG}} \)**

Phenanthrene exhibited a positive voltage dependence of WT \( I_{\text{hERG}} \) inhibition (the extent of observed inhibition increased steeply over the range of voltage-dependent activation of the channel), in common with a number of gated-state dependent small molecule inhibitors of hERG that we have studied previously (e.g. [24, 35, 42, 64, 65]). An inverse voltage dependence of \( I_{\text{hERG}} \) reduction is rarely observed, for example with protons [66, 67] or with the toxin inhibitor BeKm-1 that produces preferential closed channel inhibition [68]. Interestingly, at some voltages in the tested range, an apparent augmentation or ‘facilitation’ of \( I_{\text{hERG}} \) by phenanthrene was observed. The issue of \( I_{\text{hERG}} \) facilitation by several inhibitors has been investigated previously by Kurachi and colleagues [69–71]. These authors attributed \( I_{\text{hERG}} \) facilitation at low voltages by amiodarone, carvedilol, nifekalant and quinidine to hyperpolarization of the voltage dependence of \( I_{\text{hERG}} \) activation [69]. Remarkably, the leftward shift in \( I_{\text{hERG}} \) activation \( V_{0.5} \) by amiodarone, carvedilol, nifekalant and quinidine did not show concentration dependence for any of these compounds [69]. Our experiments have shown that phenanthrene also negatively shifts WT \( I_{\text{hERG}} \) activation (Fig. 3). Although we did not evaluate concentration dependence of the activation \( V_{0.5} \) shift for the WT channel, it is notable that we found that Y652A \( I_{\text{hERG}} \)
did not exhibit current facilitation by phenanthrene at negative voltages. This corresponded with a lack of negatively shifted activation with phenanthrene for the Y652A mutant (supplemental Fig. 4). Thus, the observed facilitation of WT $I_{hERG}$ by phenanthrene at negative test potentials can reasonably be attributed to the leftward shift in voltage dependent activation. A pharmacophore for hERG channel facilitation by inhibitors has been proposed that is comprised of one positively ionizable feature and 3 hydrophobic features [71]. Two of the facilitator molecules used in construction of this pharmacophore were nortryptiline and imipramine, both of which are comprised of 3 rings fused together with a side chain [71]. Phenanthrene is chemically simpler than either of these 3 ringed molecules, possessing 3 fused rings, but lacking an ionizable feature/size chain. Our results therefore indicate, at least in the case of phenanthrene, that 3 hydrophobic features alone can suffice for $I_{hERG}$ facilitation to occur. It has been recently suggested that facilitation of native $I_{Kr}$ by some hERG channel inhibitors may suppress proarrhythmic early afterdepolarizations [70]. Whether or not this might be the case for phenanthrene would require direct study using ventricular myocytes from an appropriate mammalian species. In zebrafish myocytes, however, the combination of the compound’s ability to inhibit $I_{Kr}$ and to accelerate deactivation led to a reduction in the native channel’s ability to generate protective outward $I_{Kr}$ transients in response to premature stimulation late in repolarization/early in diastole [15].

**Phenanthrene inhibition of $I_{hERG}$—notable and unusual features.**

The acceleration of WT $I_{hERG1a}$ and $I_{hERG}$ deactivation seen here by phenanthrene is unusual, although it is in agreement with recent observations on the effect of phenanthrene on zebrafish native $I_{Kr}$ [15]. Acceleration of $I_{hERG}$ deactivation precludes a “foot-in-door” inhibitory mechanism in which a compound’s presence in the inner cavity slows closure of the activation gate. The relationship between deactivation acceleration by phenanthrene and the compound’s ability to block the channel appears complex: both N588K and S624A $I_{hERG}$, which were largely resistant to phenanthrene inhibition, showed little or no effect of the compound on deactivation parameters (Supplemental Fig. 5), consistent with a link between current blocking and deactivation accelerating actions. On the other hand, F557L current was augmented not inhibited by phenanthrene and yet exhibited faster deactivation than in control (Fig. 5 and online supplement text), consistent with independence between current blocking and deactivation accelerating actions. It is not straightforward to unify these observations, though the F557L result indicates that deactivation acceleration is not inextricably linked to channel block by phenanthrene. There was, however, a strong correlation between the current’s deactivation rate in control solution and the derived $IC_{50}$ for phenanthrene inhibition (shown by plotting $IC_{50}$ against fast and slow inactivation $\tau$ for hERG1a, hERG1a/b and zERG; see Supplemental Fig. 6). $I_{hERG1b}$ is identical to hERG1a except for its abbreviated N terminus, which lacks the EAG domain that is known to be important for slow deactivation (Supplemental Fig. 6 and [72]). The faster deactivation of $I_{hERG1a/b}$ in control conditions was not further accelerated by phenanthrene, suggesting that a full complement of intact N-termini was not required for phenanthrene to inhibit hERG channels. However, it seems likely that conformational changes associated with fast deactivation may be involved in optimising the compound’s interaction with or access to binding residues in the pore.

Perhaps the most surprising results of this study are those pertaining to the likely interaction site through which phenanthrene inhibits $I_{hERG}$. Virtually all compounds that have been hitherto investigated have shown a strong dependence on interactions with one or both of S6 Y652 and F656: mutations of these residues typically reduce inhibitory potency [18, 50]. Indeed, structurally related halofantrine conforms to this observation [62]. In contrast, for phenanthrene, mutations at F656 failed significantly to reduce $I_{hERG}$ inhibition and the Y652A mutation increased blocking potency. Although Y652A has previously been associated with a modest (+33 mV) positive shift in the voltage dependence inactivation [73], the extent of $I_{hERG}$ inactivation at +20 mV (the command voltage used in our standard $I_{hERG}$ protocol) was shown to be similar between WT and Y652A $I_{hERG}$ [73]. Consequently, differences between WT and Y652A inactivation would not have been a significant factor in experiments investigating phenanthrene with this protocol. To our knowledge there is only one prior report of a compound for which Y652A increases inhibitory potency: capsaicin [74]. In that study, the $IC_{50}$ for capsaicin was four-fold lower for Y652A than for WT $I_{hERG}$ and the F656A mutation did not alter potency [74]. Obligatory determinants of capsaicin binding were not identified [74]. There is little structural similarity between capsaicin and phenanthrene and binding determinants are unlikely to be similar for the two compounds. The recent cryo-EM structure of hERG contains hydrophobic pockets surrounding the central cavity that are large enough to accommodate drug molecules [33]. F557 on the S5 helix lies deep within the pockets and has been implicated in the binding of a number of drugs [29, 35, 53, 75]. The recent work has implicated both F557 and M561 in hERG block by the bradycardic agent ivabradine, with mutation of M651 influencing state-dependent dynamics of aromatic residue cassettes at the pore–lipid interface [75]. These observations are consistent with the recent identification with other K+ channels that hydrophobic pockets can provide inhibitor binding sites away from the K+ permeation path [76–78].
Docking of phenanthrene identifies a site within the side pockets that can accommodate phenanthrene in structures in which the F656 side chain is rotated towards the pore. In other words, the reconfiguration of F656 into a pore-facing orientation creates space for phenanthrene to bind into a site in which interactions with F557 and M651 side chains are made (Fig. 6), consistent with our mutagenesis results (Fig. 5). Recent simulations starting with the cryo-EM structure have indicated that rotation of the F656 side chain towards the pore occurs readily and is required for optimal matching of computational analysis of drug binding with experimental mutagenesis data [29, 75, 79, 80], although in a recent bound structure for astemizole, direct interactions of the drug with F656 made only a minor contribution to overall binding [81]. Our results suggest that rotation of F656 side chains towards the pore may be required both for binding of some classical pore blockers (to maximise pore blocker interactions with aromatic side chains within the pore) and for inhibitors that bind deep within hERG pore domain side pockets (to free space for inhibitor binding). In any case, binding modes that accommodate phenanthrene deep within a pocket on the S5-S6 interface (Fig. 6) are also consistent with a minimal effect of F656 mutants on phenanthrene block (Fig. 5), in contrast with a general requirement for F656 as a binding determinant for positively charged aromatic side chains towards the pore. In other words, the reconfiguration of F656 into a pore-facing orientation creates space for phenanthrene to bind into a single hERG pore subunit is sufficient for channel block to occur. Although hydrophobic pocket binding has been identified for several K+ channel inhibitors [76–78], the mechanism by which this leads to channel inhibition remains unclear. One possibility is that inhibition involves perturbation of the selectivity filter. A curious observation is that some hERG activators which act by attenuating inactivation also bind within hERG pockets and so there may be subtle differences in blocker and activator binding that affect structure around the selectivity filter with opposite effects. Consistent with this idea a recent study on hERG activators that are likely to bind within pockets has shown that subtle chemical modification of activators can turn them into blockers [85].

In conclusion, this study demonstrates that the ubiquitous hydrocarbon pollutant phenanthrene produces an acute pharmacological block of the hERG potassium channel at concentrations compatible with tissue accumulation. We have also identified phenanthrene as a direct channel inhibitor that is mechanistically distinct from canonical hERG blockers, suggesting a broader sensitivity of this channel to adverse chemical impact than simply an off-target sensitivity to pharmaceuticals. Importantly, whilst candidate pharmaceuticals with hERG-blocking activity can be withdrawn from development, this is not the case for phenanthrene—a pervasive pollutant in air and water. As IKr is critical for human ventricular repolarization, further work is warranted to determine consequences of this effect for human ventricular electrophysiology and arrhythmia risk and whether tricyclic pollutants contribute to the morbidity and mortality associated with urban air pollution. 

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Availability of data and materials  The data for the study are included in the manuscript and supplementary information. Materials for the study will be made available on reasonable request.

Code availability  The cryo-EM structure used for this study is available at the RCSB Protein Data Bank, PDB code: 5VA2.

Declarations

Conflict of interest  The authors declare that they have no actual or potential competing interests.

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