ENDOGENOUS ION CHANNELS OF MAMMALIAN ELL LINES CHARACTERIZED WITH THE QPATCH

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A large number of mammalian cell lines are comzed the ion channel types that they endogenously

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express. Specificly we have Suitability for automated patch clamp studies

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('patchability')
Background ionic currents that may interfere Possible use for characterizing ion channels of with currents of experimentally expressed ion

interest without the need to experimentally

number of voltage and ligand gated ion channels of potential interest for the pharmaceutical industry

are endogenously expressed in several CNS and

employment of the cell lines in QPatch characteriof simple standard operation procedures (SOPs) for The tests have led to the development of a number

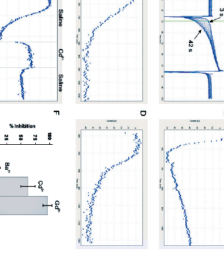
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nels, inward rectifier K * channels, acid-sensing ion channels (ASIC) and muscarinic alpha-adrenergic receptors mAChR). We have explored the applica-

channels, Ca ²⁺-release activated Ca ²⁺ (CRAC) chan non-CNS cell lines including TTX-sensitive Na

with Sophions QPatch TM automated patch clamp systems (QPatch 16 and QPatch HT) and character-

American Type Culture Collection (ATTC) for use



CELL LINE: RBL-2H3

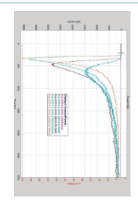
charmel currents (I conc.) in RBL-2H3 rat basophilic leukemia cells.

A: Family 14 of superimposed I conc. Figure 5.
Ca *-release activated Ca channel currents (1 GRC) in obtained after stimulation with 10 μMIns(1,4,5)P ₃ and 10 mM BAPTA. Time constant for current trace #1 and 14 are indicated. B: Current amplitude at -80 mV for 224 current

traces including those shown in panel A. C: I-t plot showing the activation of I CRAC by 15 mM BAPTA.

D: Activation of I CRAC in the absence D: Activation of CRAC in the absence of external Na * (NMDG substitution).

E: 1-t plot showing inhibition of CRAC by CdCl 2. F: Efficacy of BaCl 2. CdCl 2 and GdCl 3 in blocking I CRAC (Adapted 13:638-643, 2008).



CELL LINE: TE-671

NIELS J. WILLUMSEN

Raw current traces for the activation of mAChR currents of TE-671 human medullablastoma cells by 1-1000 μΜ

acetylcholine (Ach).





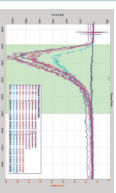
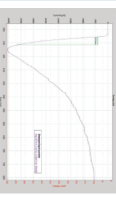






Figure 9.
Rise time of mAChR current with 1 mM of acethylcholine.









Cell line	Origin	ATTC #		Basic patchability		Na⁺ current	rrent	K + current	rent	Ca ²⁺ current	ent	LGIC current	nt
			R_{seal}	Rwholecell	Wholecells	Expression	l peak	Expression	l peak	Expression	l peak	Expression	l peak
			(GOhm)	(GOhm)	(%)	(%)	(pA)	(%)	(pA)	(%)	(pA)	(%)	(pA)
SH-SY5Y	SH-SY5Y Human neuroblastoma	CRL-2266 ™	3.42±1.01	2.08±1.01	38	28	-409±101	16 1	1040±358	ω -	-180	'	1
						(∏X insensitive)		(K _v 3.x)					
PC-12	Rat adrenal gland	CRL-1721 ™	1.22±0.35	6.47±2.30	46	-		53	263±164		1	1	1
IMR-32	Human neuroblastoma	CCL-127 ™	6.36±2.26	4.31±1.75	33	74	-377±68	21	113±13	-	-	-	-
RBL-2H3	Rat basophilic leukemia	CRL-2256 ™	1.60±0.21	1.98±0.46	77	-	1	100	-1087±79 ¹	95	-32±3	1	1
								Х _{ir}		CRAC			
TE671	Human medullablastoma	CRL-8805 ™	0.87±0.16	0.87±0.13	85	100	1525±330	•	•	•	1	100 1190±149	0±149
												mAChR ²	

Basic Properties

- The table collects the following types of data for the five cell lines that were characterized:

 1. patchability with QPatch "" based on gigaseal and whole-cell resistances and whole-cell success rate
- expression level of the endogenous ion channel species as percentage of cell recordings that exhibited the specific ionic curre whole-cell current amplitudes of the identified ion channels. The nature of the ion channels were based on use of block-

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 $Ion \, channels \, listed \, in \, parenthesis \, indicate \, most \, likely \, identity.$

The patchability was tested with physiological Ringer's solutions and without specific optimization of test conditions. For all cell lines: n>40. Where the identity of ion channels is positively known the name is given without parenthesis.

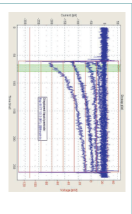
ln response to 140 mM K
 ln response to 370 μM ACh

* at-40 mV

ers, activators and in substitutions. The functional properties are presented in the figures below the properties are presented in the properties are presented in the properties are properties and the properties are properties and the properties are presented in the properties are properties and the properties are properties at the properties are properties and the properties are properties at the properties at the properties are properties at the properties at the properties are properties at the properties at

CELL LINE:SH-SY5Y

neuroblastoma cells in response to two different pulse stimulations: 1. A large outward TEA sensitive K and 2. a inward fast inactivating TTX blockable Na current followed by a slowly inactivating Cd are channel current Representative raw current traces from SH-SY5Y human 2+ sensitive + channel ⁺ current,



CELL LINE: IMR-32

Raw Ca ²⁺ (or Na ⁺) current traces in response to increasingly depolarizing test pulses. Test pulses were 200 ms long stepping from a holding potential of -110 mV to -130 mV increasing to 50 mV. The Ca ²⁺ currents were not seen in the initial experiments but were seen after the cells had been

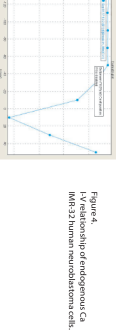


Figure 2.

Raw traces of slowly activating K currents in PC-12 rat adrenal gland cells in response to increasing depolarizing test pulses. The cell was held at -80 mV between the 50 ms test pulses that ranged from -80 to 80 mV.

CELL LINE: PC-12

SUMMARY:

2+ (or Na +) channels in

channels. It is concluded that the selection of a proper cell line in several cases may eliminate the need with the QPatch technology. It was found that each cell line exhibited a specific and characteristic expression of ion channels including sodium, potassium and calcium channels as well as ligand-gated cation and for their expression of endogenous ion channels. All cell lines proved to be well suited for exploration for expression of specific ion chann Five commercially available cell lines were tested for use with QPatch ™ automated patch clamp systems

Standard SOPs for assays based on mercially available cell lines are available from Sophion Bioscience. el genes in drug testing assays. recordings of ion channel currents of the five tested, com-

