

Gildas LOUSSOUARN

l'institut du thorax - Inserm UMR1087 - CNRS UMR6291
IRT UN - 8 quai Moncoustu
BP 70721
44007 NANTES Cedex 1
Tél. : (33) 02 28 08 01 50
E-mail : gildas.loussouarn@inserm.fr

DR2-CNRS
Marié, 2 enfants
51 ans

Cursus scientifique

- | | |
|--|---------------------|
| Chercheur invité , Université de Californie à San Francisco
Focus: collaboration avec le laboratoire de Daniel Minor sur la structure des canaux sodiques | 2012-2013 |
| CNRS –Directeur de Recherche 2^{ème} classe
Biophysique des canaux potassiques cardiaques.
INSERM UMR1087 CNRS UMR 6291. Nantes.
Directeur: R. Redon. | 2011-actuel |
| HDR - Physiologie moléculaire des canaux potassiques | Juillet 2005 |
| CNRS – Nommé Chargé de Recherche
Biophysique des canaux potassiques cardiaques.
Unité INSERM U533/UMR915. Nantes. Directeurs: Prof. D. Escande./P Pacaud | 2001-2011 |
| POST DOCTORAT
Structure et fonctions des canaux potassiques rectifiant entrant
Département de biologie cellulaire. Washington University of St Louis, USA
Directeur: Dr. C.G. Nichols. | 1997-2001 |
| DOCTORAT de l'université Paris XI
Rôle physiopathologique des canaux potassiques
INSERM CJF 96.01. Nantes. Directeur: Prof. D. Escande. | 1994-1997 |
| DEA 'Physiologie animale' de l'université de Rennes I
Physiologie de la protéine CFTR rénale
Commissariat à l'Énergie Atomique, Paris. Directeur: Dr P. Poujeol. | 1992-1993 |
| DIPLOME D'INGENIEUR AGRONOME
Ecole Nationale Supérieure Agronomique, Rennes | 1990-1992 |

Financements

- ANR Jeunes chercheuses et Jeunes chercheurs 2005 - 2008 (porteur de projet): 150 k€
- AFM 2009-2010 (porteur de projet) : 26 k€,
- Génavie 2011 (porteur de projet) : 10 k€
- Dotation FFC/SFC 2011-2012 (porteur de projet) : 80 k€

- Marie Curie 2012-2014 : 158 k€
- AFM 2013-2015 (porteur de projet) : 100 k€
- PHC-Kolmogorov 2015-2017 (porteur de projet) : 15 k€
- Génavie 2016 (porteur de projet) : 12 k€
- Fédération Française de Cardiologie (2019-2022): 300 k€
- Pari Scientifique (2019-2021): 150 k€
- ANR flash covid (2020-2021): 200 k€

Activité d'encadrement

- Encadrement de post-doctorants en France :
Kyu-Ho Park (2003-2004, maintenant post-doc à l'institut Pasteur Korea), Nicolas Rodriguez (2006-2008, maintenant Maître de Conférence, Sorbonne Université), Zeineb Es Salah Lamoureux (2011-2015, maintenant coordinatrice d'études cliniques au CHU de Nantes), Olfat Malak (2017-2018, maintenant post-doc au Buck Institute for Research on Aging, Novato, USA), Barbara Ribeiro (2018-), Jérôme Monnach (2019-).
- Encadrement d'étudiants en thèse en France et aux USA:
USA : Revell Phillips (1999-2000, maintenant Science and Technology Manager, Defense Threat Reduction Agency), Ricard Masia (2000-2001, maintenant Anatomopathologiste Massachusetts General Hospital)
France : Aziza El Harchi (2003-2006, maintenant Post-doc à l'université de Bristol), Julien Piron (2005-2008, maintenant entrepreneur), Frank Choveau (2006-2009, maintenant Post-doc à l'IHU Lyric, Bordeaux), Yassine Mohamed Amarouch (2008-2010, maintenant Professeur à l'Université de Sidi Mohamed Ben Abdellah, Fès), Fayal Abderemane Ali (2010-2013, maintenant post-doc à UCSF), Fabien Coyan (2011-2014, maintenant Business Analyst senior, Nantes), Olfat Malak (2014-2017 maintenant post-doc au Buck Institute for Research on Aging, Novato, USA).
- Encadrement d'étudiants L3/Master1/Master2 en Sciences Biologiques et Médicales
- Encadrement d'étudiants en Bioinformatique
Anne-Lise Moisdon (3ème année INSA Rennes), Sandrine Panel (MasterPro Bioinformatique Nantes), Pierre Olchesqui (Master1 Bio-informatique, Bio-statistique)

Autres activités

- **Associate editor** pour *Frontiers in Pharmacology*
- **Enseignement** en Master1 « Biologie et Biopathologie du Thorax » (Nantes), en Master1 « Sciences du médicament » (Nantes) et en Master2 « Biologie Moléculaire et Cellulaire » (Rennes)
 - **Référent** pour 'Frontiers in Cellular Neuroscience', 'Nature Communication', 'Journal of Molecular and Cellular Cardiology', 'Journal of Physiology', 'Pflugers Archiv-European Journal of Physiology', 'Journal of General Physiology', 'Journal of Biological Chemistry', 'Human Genetics', 'British Journal of Pharmacology', 'Plos One', 'Biophysical Journal', 'Expert Reviews in Molecular Medicine', 'Nature Chemical Biology', 'Cellular Physiology and Biochemistry', 'Frontiers in Pharmacology', 'Journal of Medical and Biological Engineering', 'BBA-Biomembranes'.
- **Rapporteur des thèses de :**

- Julien Dupuis ('Interaction directe récepteur-canal K_{ATP} ', 2008, CEA, Grenoble),
 - Nathalie Helix-Nieslen ('Trafficking and intracellular regulation of $K_v7.1$ potassium channel in the heart', 2008. Université de Copenhague, Danemark),
 - Reda Assal ('Méthodes de production et étude électrophysiologique de canaux ioniques : application à la pannexine1 humaine et au canal mécanosensible bactérien', 2011, Paris XI)
 - Katarzyna Niescierowicz ('Development of the ion channel-coupled receptor technology in structure function studies of G protein-coupled receptors and $kir6.2$ channel', 2013, CEA, Grenoble).
 - Camille Barbier ('Régulation de l'expression membranaire et dynamique du canal potassique $K_v1.5$ dans les cardiomyocytes atriaux', 2016, UPMC, Paris VI)
 - Catalina Reyes ('Etudes Moléculaires du canal potassique sensible à l'ATP: "gating", pathologie et optogénétique', 2016, CEA, Grenoble).
- **Examineur des thèses de :**
 - Mohamed Yassine Amarouch ('Mort subite et complexes moléculaires des canaux ioniques cardiaques', 2010, Nantes Atlantique Université),
 - Lucie Delmotte ('Fonction et modulation des canaux K_v : étude par simulation de dynamique moléculaire', 2011, Université H. Poincaré, Nancy)
 - Audrey Deyawe Kongmeneck ('investigation des mécanismes d'activation et de couplage du canal potassique voltage-dépendant $K_v7.1$ dans les cardiomyocytes à l'aide de méthodes computationnelles', 2020, Université de Lorraine, Nancy)
- **Rapporteur de l'HDR de :**
 - Delphine Bichet ('Assemblage et Fonctionnement Biophysique des Canaux Potassiques', 2016, IPMC, Valbonne).
- **Expertise** de dossiers de candidature pour le **Human Frontiers Science Program**, le **Wellcome Trust**, la **National Science Foundation**, l'**Israel Science Foundation**, le **Biotechnology and Biological Sciences Research Council**, la **Binational Science Foundation**, l'**ANR**, le **FONDECYT**, l'**Académie des Sciences d'Autriche**, l'**AFM**, la **région Bretagne**, la **région île de France**, l'**université Grenoble Alpes**, le **Musculoskeletal Research Center à St Louis**.
 - Rédaction d'un Blog pour la Société Américaine de Biophysique sur la protéine du HIV Tat. 1er Décembre 2016. <https://biophysicalsociety.wordpress.com/tag/hiv-tat/>
 - Participation à la nuit blanche des chercheurs de Nantes le 9 février 2017.
 - Membre du comité Scientifique du colloque « Microscopie électronique pour Les Sciences du Vivant à l'Université de Nantes » le 21 Juin 2019

Sociétés savantes

- *Membre de la Biophysical Society (depuis 1998)* et de la Société Française de Cardiologie

Articles originaux

43. MONTNACH J, BARÓ I, CHARPENTIER, DE WAARD M, LOUSSOUARN G, Modeling sudden cardiac death risks factors in covid-19 patients – the hydroxychloroquine and azithromycin case. *Europace* (sous presse).
42. MONTNACH J, LORENZINI M, LESAGE A, SIMON A, , NICOLAS S, MOREAU E, Céline MARIONNEAU C, BARÓ I, DE WAARD M, LOUSSOUARN G, Computer modeling of whole-cell voltage-clamp analyses to delineate guidelines for good practice of manual and automated patch-clamp. *Scientific Reports* 2021 11(1):328.
41. DE WAARD S, MONTNACH J, RIBEIRO B, NICOLAS S, FOREST V, CHARPENTIER F, MANGONI ME, GABORIT N, RONJAT M, **LOUSSOUARN G**, LEMARCHAND P, DE WAARD M. Functional Impact of BeKm-1, a High-Affinity hERG Blocker, on Cardiomyocytes Derived from Human-Induced Pluripotent Stem Cells. *Int J Mol Sci.* 2020;21:7167
40. AL SAYED ZR, CANAC R, CIMAROSTI B, BONNARD C, GOURRAUD JB, HAMAMY H, KAYSERILI H, GIRARDEAU A, JOUNI M, JACOB N, GAIGNERIE A, CHARIAU C, DAVID L, FOREST V, MARIONNEAU C, CHARPENTIER F, **LOUSSOUARN G**, LAMIRAULT G, REVERSADE B, ZIBARA K, LEMARCHAND P, GABORIT N. Human model of IRX5 mutations reveals key role for this transcription factor in ventricular conduction. *Cardiovasc Res.* 2020:cvaa259
39. MALAK OA, ABDEREMANE-ALI F, WEI Y, COYAN FC, PONTUS G, SHAYA D, MARIONNEAU C, **LOUSSOUARN G**. Up-regulation of voltage-gated sodium channels by peptides mimicking S4-S5 linkers reveals a variation of the ligand-receptor mechanism. *Sci Rep.* 2020: 10(1):5852.
38. MALAK OA, GLUHOV GS, GRIZEL AV, KUDRYASHOVA KS, SOKOLOVA OS, **LOUSSOUARN G**. Voltage-dependent activation in EAG channels follows a ligand-receptor rather than a mechanical-lever mechanism. *J Biol Chem.* 294:6506-21, 2019
37. KARLOVA M, VOSKOBOYNIKOVA N, GLUHOV GS, ABRAMOCHKIN D, MALAK OA, MULKIDJANIAN A, **LOUSSOUARN G**, SHAITAN K, STEINHOFF HJ, SOKOLOVA OS. Detergent-free solubilization of human Kv channels, expressed in mammalian cells. *Chemistry and Physics of Lipids.* 2019:50-57, 2019.
36. MONTNACH J, CHIZELLE FF, BELBACHIR N, CASTRO C, LI L, **LOUSSOUARN G**, TOUMANIANTZ G, CARCOUËT A, MEINZINGER AJ, SHMERLING D, BENITAH JP, GÓMEZ AM, CHARPENTIER F, BARÓ I. Arrhythmias precede cardiomyopathy and remodeling of Ca²⁺ handling proteins in a novel model of long QT syndrome. *J Mol Cell Cardiol*, 123:13-25, 2018.
35. BUREL S, COYAN FC, LORENZINI M, MEYER MR, LICHTI CF, BROWN JH, **LOUSSOUARN G**, CHARPENTIER F, NERBONNE JM, TOWNSEND RR, MAIER LS, MARIONNEAU C. C-terminal phosphorylation of Nav1.5 impairs FGF13-dependent regulation of channel inactivation. *J Biol Chem.* 292:17431-48, 2017.
34. MALAK OA, ES-SALAH-LAMOUREUX Z, **LOUSSOUARN G**. hERG S4-S5 linker acts as a voltage-dependent ligand that binds to the activation gate and locks it in a closed state. *Sci Rep.* 2017 Dec;7:113
33. ES-SALAH-LAMOUREUX Z, JOUNI M, MALAK OA, BELBACHIR N, AL SAYED ZR, GANDON-RENRARD M, LAMIRAULT G, GAUTHIER C, BARO I, CHARPENTIER F, ZIBARA K, LEMARCHAND P, BEAUMELLE B, GABORIT N, **LOUSSOUARN G**. HIV-Tat induces a decrease in i_{KR} and i_{KS} via reduction in phosphatidylinositol-(4,5)-bisphosphate availability. *J Mol Cell Cardiol.* 99:1-13, 2016.
32. PORTERO V, LE SCOUARNEC S, ES-SALAH-LAMOUREUX Z, BUREL S, GOURRAUD JB, BONNAUD S, LINDENBAUM P, SIMONET F, VIOLLEAU J, BARON E, MOREAU E, SCOTT C, CHATEL S, **LOUSSOUARN G**, O'HARA T, MABO P, DINA C, LE MAREC H, SCHOTT JJ, PROBST V, BARO I, MARIONNEAU C, CHARPENTIER F, REDON R. Dysfunction of the voltage-gated k⁺ channel $\beta 2$ subunit in a familial case of Brugada syndrome. *J Am Heart Assoc.* 5(6), 2016.
31. JOUNI M, SI-TAYEB K, ES-SALAH-LAMOUREUX Z, LATYPOVA X, CHAMPON B, CAILLAUD A, RUNGOAT A, CHARPENTIER F, **LOUSSOUARN G**, BARÓ I, ZIBARA K, LEMARCHAND P, GABORIT N. Toward Personalized Medicine: Using Cardiomyocytes Differentiated From Urine-Derived Pluripotent Stem Cells to Recapitulate Electrophysiological Characteristics of Type 2 Long QT Syndrome. *J Am Heart Assoc.* 4(9), 2015. IF 4.31

30. MORENO C, DE LA CRUZLA A, OLIVERAS A, KHARCHE SR, GUIZY M, COMES N, STARÝ T, RONCHI C, ROCCHETTI M, BARÓ I, **LOUSSOUARN G**, ZAZA A, SEVERI S, FELIPE A, VALENZUELA C. Marine n-3 PUFAs modulate IKs gating, channel expression, and location in membrane microdomains. *Cardiovasc Res*. 105:223-32, 2015. IF 5.94 (IF 2014)
29. BEZIAU DM, BARC J, O'HARA T, LE GLOAN L, AMAROUCHE YM, SOLNON A, PAVIN D, LECOINTE S, BOUILLET P, GOURRAUD JB, GUICHENEY P, DENJOY I, REDON R, MABO P, LE MAREC H, **LOUSSOUARN G**, KYNDT F, SCHOTT JJ, PROBST V, BARÓ I. Complex Brugada syndrome inheritance in a family harbouring compound SCN5A and CACNA1C mutations. *Basic Res Cardiol*. 109:446, 2014, IF 5.96.
28. COYAN F, ABDEREMANE-ALI F, AMAROUCHE MY, PIRON J, MORDEL J, NICOLAS CS, STEENMAN M, MEROT J, MARIONNEAU C, THOMAS A, BRASSEUR R, BARÓ I, **LOUSSOUARN G**. A Long QT Mutation Substitutes Cholesterol for Phosphatidylinositol-4,5-Bisphosphate in KCNQ1 Channel Regulation. *PLoS One*. 9:e93255, 2014. IF 3.73
27. SHAYA D, FINDEISEN F, ABDEREMANE-ALI F, ARRIGONI C, WONG S, NURVA SR, **LOUSSOUARN G**, MINOR DL, Jr. Structure of a Prokaryotic Sodium Channel Pore Reveals Essential Gating Elements and an Outer Ion Binding Site Common to Eukaryotic Channels. *J Mol Biol*. 426:467-83, 2014. IF 2.48
26. ABDEREMANE-ALI F, ES-SALAH-LAMOUEUX Z, DELEMOTTE L, KASIMOVA MA, LABRO AJ, SNYDERS DJ, FEDIDA D, TAREK M., BARÓ I, **LOUSSOUARN G**. Dual effect of Phosphatidylinositol-4,5-Bisphosphate on Shaker K⁺ channels. *J Biol Chem* 287:36158-67, 2012. IF 4.65
25. LAURENT G., SAAL S., AMAROUCHE M.Y., BEZIAU D.M., MARSMAN R.F.J., FAIVRE L., BARC J., DINA C., BERTAUX G., BARTHEZ O., THAUVIN-ROBINET C., CHARRON P., FRESSART V., MALTRET A., VILLAIN E., BARON E., MEROT J., TURPAULT R., COUDIERE Y., CHARPENTIER F., SCHOTT JJ, **LOUSSOUARN G**., WILDE A., WOLF JE., BARÓ I, KYNDT F., PROBST V. F., Multifocal Ectopic Purkinje-related Premature Contractions: a new SCN5A-related cardiac channelopathy. *J Am Coll Cardiol* 60, 144-156, 2012. IF 14.29
24. LABRO AJ, BOULET IR, CHOVEAU F, MAYEUR E, BRUYNS T, **LOUSSOUARN G**, RAES AL, SNYDERS DJ. The S4-S5 linker of KCNQ1 channels forms a structural scaffold with the S6 segment controlling gate closure. *J Biol Chem*. 286:717-25, 2011. IF 5.33
23. CHOVEAU F, RODRIGUEZ N, ABDEREMANE ALI F, LABRO AJ, ROSE T, DAHIMENE S, BOUDIN H, LE HENAFF C, ESCANDE D, SNYDERS DJ, CHARPENTIER F, MEROT J, BARO I, **LOUSSOUARN G**. KCNQ1 channels voltage dependency through a voltage-dependent binding of the S4-S5 linker to the pore domain. *J Biol Chem*. 286:707-16, 2011. IF 5.33
22. PIRON J, CHOVEAU FS, AMAROUCHE MY, RODRIGUEZ N, CHARPENTIER F, MEROT J, BARO I, **LOUSSOUARN G**. KCNE1-KCNQ1 osmoregulation by interaction of phosphatidylinositol-4,5-bisphosphate with Mg²⁺ and polyamines. *J Physiol*, 588:3471-3483, 2010. IF 4.76 → **Cf aussi "Comments" par N. Gamper in J Physiol. 588:3631-2, 2010**
21. RODRIGUEZ N, AMAROUCHE MY, MONTNACH J, PIRON J, LABRO AJ, CHARPENTIER F, MEROT J, BARO I, **LOUSSOUARN G**. Phosphatidylinositol-4,5-bisphosphate (PIP₂) stabilizes the open pore conformation of the Kv11.1 (hERG) channel. *Biophysical J*, 99:1110-1118, 2010. IF 4.39
20. CHOVEAU FS, EL HARCHI A, RODRIGUEZ N, LOUERAT-ORIOU B, BARÓ I, DEMOLOMBE S, CHARPENTIER F, **LOUSSOUARN G**. Transfer of rolf S3-S4 linker to hERG eliminates activation gating but spares inactivation. *Biophysical J*, 97:1323-1334, 2009. IF 4.39
19. HAISSAGUERRE M, CHATEL S, SACHER F, WEERASOORIYA R, PROBST V, **LOUSSOUARN G**, HORLITZ M, LIERSCH R, SCHULZE-BAHR E, WILDE A, KAAB S, KOSTER J, RUDY Y, LE MAREC H, SCHOTT JJ. Ventricular fibrillation with prominent early repolarization associated with a rare variant of KCNJ8/KATP channel. *J Cardiovasc Electrophysiol* 20:93-8, 2009. IF 3.70
18. PEROZ D, DAHIMENE D, BARÓ I, **LOUSSOUARN G**, MEROT J. LQT1 associated mutations increase KCNQ1 proteasomal degradation independently of derlin-1. *J Biol Chem*. 284:5250-6, 2009. IF 5.33
17. NICOLAS C, PARK KH, EL HARCHI A, CAMONIS J, KASS RS, ESCANDE D, MEROT J, **LOUSSOUARN G**, LE BOUFFANT F, BARÓ I. IKs response to protein kinase A-dependent KCNQ1 phosphorylation requires direct interaction with microtubules. *Cardiovasc Res* 79:427-35, 2008. IF 5.95

16. BAUDOIN SJ, ANGIBAUD J, **LOUSSOUARN G**, BONNAMAIN V, MATSUURA A, KINEBUCHI M, NAVEILHAN P, BOUDIN H. The signaling adaptor protein CD3zeta is a negative regulator of dendrite development in young neurons. *Mol Biol Cell*. 19:2444-2456, 2008. IF 5.56
15. **LOUSSOUARN G**, MARTON J, NICHOLS CG. Molecular Basis of Inward-rectification: Structural features of the blocker defined by extended polyamine analogs. *Molecular Pharmacology* **68**, 298-304, 2005. IF 4.61
14. PARK KH, PIRON J, DAHIMENE D, MÉROT J, BARÓ I, ESCANDE D, **LOUSSOUARN G**. Impaired KCNQ1/KCNE1 and Phosphatidylinositol-4,5-bisphosphate interaction underlies the Long QT Syndrome. *Circulation Research* **96**, 730-739, 2005. IF 9.41
13. ROYER A, DEMOLOMBE S, EL HARCHI A, LE QUANG K, PIRON J, TOUMANIANTZ G, MAZURAS D, BELLOCQ C, LANDE G, TERRENOIRE C, MOTOIKE HK, CHEVALLIER JC, **LOUSSOUARN G**, CLANCY CE, ESCANDE D, CHARPENTIER F. Expression of human ERG K(+) channels in the mouse heart exerts anti-arrhythmic activity. *Cardiovascular Research* **65**, 128-137, 2005. IF 5.28
12. KURATA H.T., PHILLIPS L.R., ROSE T., **LOUSSOUARN G**, HERLITZE S., FRITZENSCHAFT H., ENKVETCHAKUL D., NICHOLS C.G., BAUKROWITZ T. Molecular basis of inward rectification: polyamine interaction sites located by combined channel and ligand mutagenesis. *Journal of General Physiology* **124**, 541-554, 2004. IF 5.10
11. **LOUSSOUARN G**, PARK KH, BELLOCQ C, BARO I, CHAPENTIER, F, ESCANDE D. Phosphatidylinositol-4,5-bisphosphate, PIP2, Controls KCNQ1/KCNE1 Voltage-gated Potassium Channels: a functional homology between voltage-gated and inward rectifier K⁺ channels. *EMBO J*. **22**, 5412-21, 2003. IF 10.45
10. **LOUSSOUARN G**, PIKE LJ, ASHCROFT FM, MAKHINA EN, NICHOLS CG. Dynamic sensitivity of ATP-sensitive K⁺ channels to ATP. *Journal of Biological Chemistry* **276**, 29098-29103, 2001. IF 7.26
9. **LOUSSOUARN G**, ROSE T, PHILLIPS R, MASIA R, NICHOLS CG. Flexibility of the Kir6.2 inward rectifier channel pore. *Proc. Natl. Acad. Sci. USA* **98**, 4227-4232, 2001. IF 10.90
8. ENKVETCHAKUL D, **LOUSSOUARN G**, MAKHINA EN, NICHOLS CG. ATP interaction with the open state of the K_{ATP} channels. *Biophysical Journal* **80**, 719-728, 2001. IF 4.64
7. GRIBBLE FM., **LOUSSOUARN G**, TUCKER SJ, ZHAO C, NICHOLS C, ASHCROFT FM. A novel method for measurement of sub-membrane ATP concentration. *Journal of Biological Chemistry* **278**, 30046-30049, 2000. IF 7.26
6. ENKVETCHAKUL D, **LOUSSOUARN G**, SHYNG SL, MAKHINA EN, NICHOLS CG. A critical gating transition in K_{ATP} channels: ATP stabilizes a closed-state. *Biophysical Journal* **78**, 2334-2348, 2000. IF 4.63
5. **LOUSSOUARN G**, MAKHINA EN, ROSE T, NICHOLS CG. Structure and dynamics of the pore of inward rectifier K_{ATP} channels. *Journal of Biological Chemistry* **275**, 1137-1144, 2000. IF 7.26
4. POLLARD H, REMY JS, **LOUSSOUARN G**, DEMOLOMBE S, BEHR JP, ESCANDE D. Polyethylenimine but not cationic lipids promotes transgene nuclear targeting in mammalian cells. *Journal of Biological Chemistry* **273**: 7507-7511, 1998. IF 7.20
3. MOHAMMAD-PANAH R, DEMOLOMBE S, RIOCHET D, LEBLAIS V, **LOUSSOUARN G**, POLLARD H, BARÓ I, ESCANDE D. Hyperexpression of recombinant CFTR in heterologous cells alters its physiological properties. *American Journal of Physiology* **274**: C310-C318, 1998. IF 3.08
2. **LOUSSOUARN G**, CHARPENTIER F, MOHAMMAD-PANAH R, KUNZELMANN K, BARO I, ESCANDE D. KvLQT1 potassium channel but not Isk is the molecular target for chromanol 293B. *Molecular Pharmacology* **52**: 1131-1136, 1997. IF 4.92
1. **LOUSSOUARN G**, DEMOLOMBE S, MOHAMMAD-PANAH R, ESCANDE D, BARÓ I. Expression of CFTR controls cAMP-dependent activation of epithelial K⁺ currents. *American Journal of Physiology*, **271**: C1565-1573, 1996. IF 3.08

Revues

11. BEAUMELLE B, TOTH P, MALAK OA, CHOPARD C, **LOUSSOUARN G**, VITALE N. Phosphatidylinositol (4,5)-bisphosphate-mediated pathophysiological effect of HIV-1 Tat protein. *Biochimie* 141:80,2017.
10. **LOUSSOUARN G**, STERNBERG D, NICOLE S, MARIONNEAU C, LE BOUFFANT F, TOUMANIANTZ G, BARC J, MALAK O, FRESSART V, PEREON Y, BARO I AND CHARPENTIER F. Physiological and pathophysiological insights of Nav1.4 and Nav1.5 comparison. *Front. Pharmacol.* 6:314, 2015.
9. **LOUSSOUARN G**. Les courants potassiques rectifiants entrants cardiaques. *Archives de Maladies du cœur et des Vaisseaux Pratique.* 237 :14-27, 2015.
8. ABRIEL H, DE LANGE E, KUCERA JP, **LOUSSOUARN G**, TAREK M. Computational tools to investigate genetic cardiac channelopathies. *Front Physiol.* 2013, 4:39. IF 3.5
7. COYAN FC, **LOUSSOUARN G**. Cholesterol regulation of ion channels: Crosstalk in proteins, crosstalk in lipids. *Channels (Austin).* 7(6), 2013. IF 2.16
6. **LOUSSOUARN G**, TAREK M. Mechanisms of ion channels voltage-dependency: all about molecular sensors, gates, levers, locks and grease. *Front. Pharmacol.* 3:174, 2012. IF 3.8 (IF 2015)
5. CHOVEAU FS, ABDEREMANE ALI F, COYAN FC, ES-SALAH-LAMOUEUX Z, BARÓ I, **LOUSSOUARN G**. Opposite effects of the S4–S5 linker and PIP₂ on voltage-gated channel function: KCNQ1/KCNE1 and other channels. *Front. Pharmacol.* 3:125, 1-16, 2012. IF 3.8 (IF 2015)
4. **LOUSSOUARN G**, BARÓ I. Neural modulation of ion channels in cardiac arrhythmias: Clinical implications and future investigations. *Heart Rhythm.* 7:847-849, 2010. IF 4.44
3. CHARPENTIER F, MEROT J, **LOUSSOUARN G**, BARÓ I. Delayed rectifier K(+) currents and cardiac repolarization. *J Mol Cell Cardiol.* 48:37-44, 2010. IF 5.05
2. PEROZ D, RODRIGUEZ N, CHOVEAU F, BARÓ I, MEROT J, **LOUSSOUARN G**. Kv7.1 (KCNQ1) properties and channelopathies. *J Physiol.* **586**, 1785-1789, 2008. IF 4.65
1. **LOUSSOUARN G**, ROSE T, NICHOLS CG. Structural Basis of Inward Rectifying Potassium Channel Gating. *Trends in Cardiovascular Medicine* **12**, 253-258, 2002. IF 3.40

Chapitres d'ouvrage

4. DEYAWA A, KASIMOVA MA, DELEMOTTE L, **LOUSSOUARN G**, TAREK M. Studying Kv Channels Function using Computational Methods. Dans “Methods Mol Biol. Potassium Channels” eds. Show-Ling Shyng, Francis Valiyaveetil and Matt Whorton, Humana Press Methods Mol Biol. 2018; 1684:321-341.
3. NOVOSELETSKY V, MALAK OA, **LOUSSOUARN G**, SOKOLOVA OS. Building Atomic Models of the Ion Channels Based on Low Resolution Electron Microscopy Maps and Homology Modeling. Dans “Methods Mol Biol. Potassium Channels” eds. Show-Ling Shyng, Francis Valiyaveetil and Matt Whorton, Humana Press Methods Mol Biol. 2018; 1684:305-319.
2. NICOLAS C, PIRON J, RODRIGUEZ N, CHOVEAU F, DAHIMENE D, PEROZ D, MÉROT J, BARÓ I and **LOUSSOUARN G**. Molecular description of KCNE1/KCNQ1 cardiopathies. Dans “Ion Channels Biophysics and Diseases”, Research Signpost Publisher, ed. H Duclohier 2009; 13-41.
1. **LOUSSOUARN G**, BARÓ I, ESCANDE D. KCNQ1 K⁺ channel-mediated cardiac channelopathies. Dans “Methods Mol Biol. Ion Channels methods and protocols” eds. JD Stockand. et MS Shapiro, Humana Press 2006; 337:167-83.

Séminaires et Communications invitées :

18. Opposite effects of the S4–S5 linker and the phospholipid PIP₂ on voltage-gated channel function: KCNQ1, hERG and other channels. Young Investigator School in Nantes. Nantes, IRS-UN, 01-02 Juin 2017

17. hERG S4-S5 linker acts as a voltage-dependent ligand that binds to the activation gate and locks it in a closed state . Young Investigator School in Moscow. Moscou, Université Lomonosov, 26 Mai 2016
16. Molecular mechanisms of Kv and Nav voltage dependent gating: a unified model. Workshop “Computational tools to investigate genetic channelopathies”, Beatenberg, 10-12 Janvier 2016
15. Canaux sodiques musculaires : vers une nouvelle approche thérapeutique ? Hotel Quintessia, Orvault, 21 Septembre 2015
14. Mécanismes moléculaires de dépendance au potentiel des canaux potassiques et sodiques, Institut du Cerveau et de la Moelle, 14 Novembre 2014.
13. Mécanismes moléculaires de la dépendance au potentiel de Nav1.4. Groupe Résocanaux, Hôpital Salpêtrière, 20 Juin 2014.
12. Insights into Kv and Nav voltage-dependent gating Implications in channelopathies. Workshop “Computational tools to investigate genetic channelopathies”, Beatenberg, 9-12 Octobre 2013.
11. Voltage dépendance du canal KCNQ1 et son implication dans les canalopathies. Séminaire externe, Centre de Recherche en Neurobiologie et Neurophysiologie de Marseille, 6 Décembre 2010.
10. Simulating ion channels activity: From currents to action potential. Mathematical modeling and computing in electrocardiology (international meeting), Nantes, 8-9 Juin 2009.
9. Intérêt de la modélisation en électrophysiologie cardiaque. Automnales du GRRC. Fontevraud, 2-4 Décembre 2009.
8. Insights into KCNQ1 and HERG voltage-dependent gating. Implications in cardiopathies. Ion channels Symposium 2008, Copenhague, Danemark, 22-23 Mai 2008.
7. Molecular mechanism underlying KCNQ1 voltage dependency: the channel gate is locked closed by the S4-S5 linkers. Joint meeting of the Slovak Physiological Society, the Physiological Society and the Federation of European Physiological Society. Bratislava, 11-14 Septembre 2007.
6. Phosphoinositides modulation of ion channels. Biosensors International Summerschool. Berder, 25-31 Août 2007.
5. Homologies in molecular physiology between Kv and Kir channels. Colloque ANR. Nancy, 9-10 Mai 2007.
4. Homologies structurales et fonctionnelles chez les canaux potassiques. Ecole polytechnique, Paris, Janvier 2003.
3. Polyamine interaction with inward rectifier K channels: *In vitro* and *in silico* correlates. Atelier bioinformatique, Le Croisic, Novembre 2002.
2. Polyamine conformation and localization when blocking inward rectifier K channels. 13ème colloque de l’association « canaux ioniques », Giens, Septembre 2002.
1. Cinétique d’expression hétérologue de la protéine CFTR. Atelier des protéines membranaires, Vaincre La Mucoviscidose, Paris, Juin 1996.

Abstracts

57. RIBEIRO DE OLIVEIRA MENDES B, MONTNACH J, NICOLAS S, OLLIVIER B, CHATELAIN FC, FOREST V, FELICIANIELI S, KYNDT F, PROBST V, LESAGE F, DENJOY I, **LOUSSOUARN G**, DE WAARD M. Functional Characterization of KCNH2 genetic variants, encoding hERG potassium channel, as a clinically-relevant information for type 2 LQTS syndrome. Archives of Cardiovascular Diseases. 12:257, Octobre 2020.
56. MONTNACH J, RIBEIRO DE OLIVEIRA MENDES B, DE WAARD S, CORREIA E, NICOLAS S, FOREST V, JOPLING C, GABORIT N, LEMARCHAND P, **LOUSSOUARN G**, DE WAARD M. Optical control of hERG channel activity using a photosensitive Bekm-1 blocker. Archives of Cardiovascular Diseases. 12:258, Octobre 2020.
- G**, IWAMOTO T, BARC J, PROBST V, KYNDT F, REDON R, MAKITA N, CHARPENTIER F, SCHOTT JJ. Identification and functional characterization of the first SLC8A1 variants responsible for idiopathic

ventricular fibrillation, J wave syndrome and short QT syndrome. European Working Group in Cardiac Electrophysiology, Essen, Juin 2018.

54. CHIZELLE F, ISHIKAWA T, **LOUSSOUARN G**, IWAMOTO T, BARC J, PROBST V, KYNDT F, REDON R, MAKITA N, CHARPENTIER F, SCHOTT JJ. Identification and functional characterization of the first variants in the SLC8A1 gene responsible for idiopathic ventricular fibrillation. *Printemps de la Cardiologie*, Montpellier, Avril 2018.

53. MALAK OA, ES-SALAH-LAMOUREUX Z, **LOUSSOUARN G**. hERG s4-s5 linker acts as a voltage-dependent ligand that binds to the activation gate and locks it in a closed state. *Archives of Cardiovascular Diseases*. 9:203, Nantes, Avril 2017.

52. ES-SALAH-LAMOUREUX Z, JOUNI M, MALAK OA, BELBACHIR N, AL SAYED ZR, GANDON-RENARD M, LAMIRAULT G, GAUTHIER C, BARO I, CHARPENTIER F, ZIBARA K, LEMARCHAND P, BEAUMELLE B, GABORIT N, **LOUSSOUARN G**. Hiv-Tat induces a decrease in I-Kr and I-Ks via reduction in Phosphatidy-Inositol-(4,5)-Bisphosphate availability. *Biophysical J.*, 112: 405A, Nouvelle Orleans, Fev 2017

51. ES-SALAH-LAMOUREUX Z, JOUNI M, MALAK OA, BELBACHIR N, AL SAYED ZR, GANDON-RENARD M, LAMIRAULT G, GAUTHIER C, BARO I, CHARPENTIER F, ZIBARA K, LEMARCHAND P, BEAUMELLE B, GABORIT N, **LOUSSOUARN G**. A molecular substrate for long QT in HIV patients: Tat protein reduces IKr in human induced pluripotent stem cells-derived cardiomyocytes. *Channelpathy meeting*, Paris, Juin 2016.

50. ES-SALAH-LAMOUREUX Z, JOUNI M, MALAK OA, BELBACHIR N, GANDON-RENARD M, LAMIRAULT G, GAUTHIER C, BARO I, CHARPENTIER F, ZIBARA K, LEMARCHAND P, BEAUMELLE B, GABORIT N, **LOUSSOUARN G**. A molecular substrate for long qt in hiv patients: tat protein reduces ikr in human induced pluripotent stem cells-derived cardiomyocytes. *Biophysical J.*, 110: 103A, Los Angeles, Février 2016.

49. MALAK OA, ES-SALAH-LAMOUREUX Z, **LOUSSOUARN G**. hERG s4-s5 acts as a voltage-dependent ligand binding the activation gate and locking it in a closed state. *Biophysical J.*, 110: 278A, Fev 2016.

48. ES-SALAH-LAMOUREUX Z, JOUNI M, LAMIRAULT G, GAUTHIER C, BARO I, CHARPENTIER F, ZIBARA K, LEMARCHAND P, BEAUMELLE B, GABORIT N, **LOUSSOUARN G**. Mechanism of induction of cardiac arrhythmias in HIV patients. *European Working Group in Cardiac Electrophysiology*, Milan, Juin 2015.

47. JOUNI M, SI-TAYEB K, ES-SALAH-LAMOUREUX Z, LATYPOVA X, CHAMPON B, RUNGOAT A, CHARPENTIER F, **LOUSSOUARN G**, ZIBARA K, LEMARCHAND P, GABORIT N. Using cardiomyocytes differentiated from urine-derived hiPSCs to recapitulate electrophysiological characteristics of LQT2 syndrome. *Archives of Cardiovascular Diseases* 7:165, Avr 2015.

46. JOUNI M, CHAMPON B, CHARPENTIER F, LOUSSOUARN G, BARO I, RUNGOAT A, LATYPOVA X, DAVID L, DEREVIER A, ZIBARA K, LEMARCHAND P, GABORIT N. Electrophysiological characterization of a novel SCN5A mutation causing Brugada syndrome, using cardiomyocytes differentiated from hiPSCs.. *Archives of Cardiovascular Diseases* 7:169, Avr 2015.

45. FINDEISEN F, SHAYA D, ABDEREMANE-ALI F, ARRIGONI C, **LOUSSOUARN G**, MINOR DL, Jr. Identification of a Determinant of High Affinity Calcium Binding in the Selectivity Filter of a Mammalian Calcium Channel. *Biophysical J.*, 106: 331A, Fev 2014.

44. ARRIGONI C, SHAYA D, FINDEISEN F, ABDEREMANE-ALI F, **LOUSSOUARN G**, MINOR DL, Jr. Structure of a Prokaryotic Sodium Channel Pore Reveals Essential Gating Elements and an Outer Ion Binding Site Common to Eukaryotic Channels. *Biophysical J.*, 106: 130A, Fev 2014.

43. COYAN F., AMAROUCHE M.Y., ABDEREMANE-ALI F., PIRON J., MORDEL J., NICOLAS C.S., STEENMAN M., MEROT J., MARIONNEAU C., THOMAS A., BRASSEUR R., BARÓ I., **LOUSSOUARN G**. In KCNQ1 Channels, a Long QT Mutation Induces a Regulation by Phosphatidylinositol-4,5-bisphosphate in channel regulation. *Biophysical J.*, 106: 140A, Fev 2014.

42. COYAN F., AMAROUCHE M.Y., PIRON J., MORDEL J., NICOLAS C.S., MEROT J., THOMAS A., BRASSEUR R., CHARPENTIER F., BARÓ I., **LOUSSOUARN G**. A long QT mutation substitutes cholesterol for phosphatidylinositol-4,5- Bisphosphate in KCNQ1 channel regulation. *24es Journées Européennes de la Société Française de Cardiologie*, Jan 2014.

41. ABDEREMANE-ALI F, ES-SALAH-LAMOUREUX Z, DELEMOTTE L, KASIMOVA MA, LABRO AJ, SNYDERS DJ, FEDIDA D, TAREK M., BARÓ I, **LOUSSOUARN G**. Dual effect of PIP2 on Shaker K⁺ channels. *Biophysical J.*, 104:464A, 2013.

40. LAURENT G., SAAL S., AMAROUCHE M.Y., BEZIAU D.M., MARSMAN R.F.J., DINA C., CHARRON P., MALTRET A., TURPAULT R., WILDE A., WOLF JE., **LOUSSOUARN G.**, KYNDT F., PROBST V. F,

- BARÓ I. R222Q Nav1.5 mutation associated with a new SCN5A-related cardiac arrhythmia. Biophysical Society meeting 2012. *Biophysical J.*, 102:527A, 2012.
39. CHATEL S., SACHER F., CHAIGNE S., LOUSSOUARN G., BARO I., KYNDT F., LE MAREC H., PROBST V., HAISSAGUERRE M., SCHOTT JJ. KATP channels and Early Repolarization Syndrome. *European Heart Journal*, 33 :971, 2012.
38. PIRON J., AMAROUCHE M.Y., COYAN F., MORDEL J., NICOLAS C.S., THOMAS A., BRASSEUR R., CHARPENTIER F., MEROT J., BARÓ I., **LOUSSOUARN G.** KCNQ1-R539W mutation substitutes cholesterol for phosphatidylinositol-4,5-bisphosphate in channel regulation. Biophysical Society meeting 2011. *Biophysical J.*, 100:428, 2011.
37. AMAROUCHE M.Y., LAURENT G., SAAL S., FAIVRE L., BERTAUX G., FALCON-EICHER S., BARTHEZ O., THAUVIN-ROBINET C., CHARRON P., RICHARD P., DE CHILLOU C., PROBST V., BARON E., BARÓ I., BARC J, SCHOTT JJ, MEROT J., COUDIERE Y., TURPAULT R., **LOUSSOUARN G.**, KYNDT F., WOLF JE. A novel familial cardiac arrhythmia associated with a dilated cardiomyopathy is due to a mutation in SCN5A. *Printemps de la cardiologie*. 15-17 avril 2010, Nantes. *Archives of Cardiovascular diseases*. 2: 41, 2010.
36. LAURENT G, SAAL S, AMAROUCHE MY, FAIVRE L, THAUVIN-ROBINET C, CHARRON P, RICHARD P, PROBST V, BARON E, BARÓ I, BARC J, SCHOTT JJ, MEROT J, TURPAULT R, COUDIERE Y, **LOUSSOUARN G.** Kyndt F, Wolf JE. Multifocal Ectopic Purkinje Tachycardia: A New Familial Syndrome. American Heart Association Scientific Session. November 2010. *Circulation*. 122: A17090, 2010.
35. AMAROUCHE M.Y., SAAL S., BERTAUX G., FAIVRE L., FALCON-EICHER S., BARON E., THAUVIN-ROBINET C., BARC J, BARTHEZ O, MEROT J., CHARRON P., RICHARD P., PROBST V., DE CHILLOU C., BARÓ I, SCHOTT JJ, WOLF JE, KYNDT F., **LOUSSOUARN G.**, LAURENT G. Mutation of cardiac Nav1.5 in a novel autosomic dominant in a hisian-type arrhythmia, associated with dilated cardiomyopathy. Biophysical Society meeting 2010. *Biophysical J.*, 1616-Pos, 2010.
34. CHOVEAU F., EL HARCHI A., RODRIGUEZ N., LOUERAT-ORIOU B., BARÓ I., DEMOLOMBE S., CHARPENTIER F., **LOUSSOUARN G.** Transfer of rolf S3-S4 linker to hERG eliminates activation gating but spares inactivation? *GRRC*, 26^{ème} congrès du GRRC. 2-3 avril 2009, Nancy. *Archives of Cardiovascular diseases*. 102-S62, 2009.
33. EL HARCHI A, CHOVEAU FS, RODRIGUEZ N, LOUERAT-ORIOU B, BARO I, DEMOLOMBE S, CHARPENTIER F, **LOUSSOUARN G.** Transfer of rolf S3-S4 linker to hERG eliminates activation gating but spares inactivation. Biophysical Society meeting 2009. *Biophysical J.*, 977-Pos, 2009.
32. BAUDOIN S.J., ANGIBAUD J., **LOUSSOUARN G.**, NAVEILHAN P., BOUDIN H. The role of the signalling protein CD3zeta in neuronal development: from dendritic outgrowth regulation to synaptic expression. *FENS* 2008.
31. RODRIGUEZ N., BARÓ I., **LOUSSOUARN G.** Does The Phosphoinositide PIP₂ Stabilize HERG Channel Open State? Biophysical Society meeting 2008. *Biophysical J.*, 1332-Pos, 2008.
30. CHOVEAU F., DAHIMENE D., RODRIGUEZ N., LE HENAFF C., MEROT J., ESCANDE D., BARÓ I., **LOUSSOUARN G.** Des fragments peptidiques du canal potassique cardiaque KCNQ1 pour contrôler sa voltage-dépendance. *GRRC*. 24^{ème} congrès du GRRC. 25-27 avril 2007, Tours.
29. NICOLAS C.S., PARK K.H., EL HARCHI A., CAMONIS J., KASS R.S., ESCANDE D., MEROT J., **LOUSSOUARN G.**, LE BOUFFANT F., BARÓ, I. Iks response to PKA-dependent phosphorylation requires direct interaction with the microtubules. Biophysical Society meeting 2007. *Biophysical J.*, L160-Pos (late abstract), 2007.
28. PIRON J., DAHIMENE D., MEROT J., ESCANDE D., BARÓ I., **LOUSSOUARN G.** Phosphoinositide PIP₂-KCNQ1 interaction implicated in channel osmoregulation. Biophysical Society meeting 2007. *Biophysical J.*, 509-Pos, 2007.
27. CHOVEAU F., DAHIMENE D., LE HENAFF C., MEROT J., ESCANDE D., BARÓ I., **LOUSSOUARN G.** Operation of a voltage-dependent channel by its S4-S5 linker. Biophysical Society meeting 2007. *Biophysical J.*, 883-Plat, 2007.
26. KURATA H.T., PHILLIPS L.R., ROSE T., **LOUSSOUARN G.**, HERLITZE S, FRITZENSCHAFT H, ENKEVETCHAKUL D, BAUKROWITZ T, NICHOLS CG. Molecular basis of inward rectification: Localizing Polyamine Binding Sites in Kir channels. Biophysical Society meeting 2005. *Biophysical J.*, 1390-Pos, 2005.
25. PARK K.H., DAHIMENE D., MÉROT J., BARÓ I., ESCANDE D., **LOUSSOUARN G.** Reduced phosphatidylinositol-4,5-bisphosphate and KCNQ1 channel interaction in the Long QT Syndrome. Biophysical Society meeting 2005. *Biophysical J.*, 2220-Pos, 2005.

24. PARK K.H., LE BOUFFANT, F., ALCOLEA S., MEROT J., CAMONIS J., **LOUSSOUARN G.**, BARO, I. Beta-tubulin, a new regulator of the slow delayed rectifier potassium current I_{ks}. American Heart Association Scientific Session. Novembre 2004, New Orleans.
23. PARK K.H., LE BOUFFANT, F., ESCANDE, D., **LOUSSOUARN G.**, BARO, I. Beta-tubulin, a new regulator of the slow delayed rectifier potassium current I_{ks}. GRRRC. 21^{ème} congrès du GRRRC. 22-23 avril 2004, La Baule. Publié dans les Archives des maladies des vaisseaux et du cœur.
22. PARK, K.H., **LOUSSOUARN G.**, ESCANDE D. Régulation du complexe KCNQ1/KCNE1 par le PIP2 et le MgATP intracellulaire. GRRRC. 20^{ème} congrès du GRRRC. 15 et 16 avril 2003. Publié dans les Archives des maladies des vaisseaux et du cœur.
21. **LOUSSOUARN G.**, ESCANDE D. Functional KCNQ1/KCNE1 channel complex requires both intracellular PIP₂ and MgATP. Publié dans Biophysical J., 2678-Pos, 2003
20. PHILLIPS R., **LOUSSOUARN G.**, NICHOLS C.G. Cold draft through the tepee: MTSEA access to the inner vestibule of ATP closed Kir6.2 channels implies gating does not occur at the smokehole. Publié dans Biophysical J., 2885-Pos, 2002
19. **LOUSSOUARN G.**, ENKVETCHAKUL D., VALASINAS A., REDDY V.K., BASU H.S., FRYDMAN B., NICHOLS C.G. Using polyamine analogs as probes to define structural constraints for block of inward rectifier K channels. Biophysical J., 2887-Pos, 2002.
18. ENKVETCHAKUL D., **LOUSSOUARN G.**, MAKHINA E.N., NICHOLS C.G. ATP interaction with the open state of K_{ATP} channels: evidence for allosteric gating mechanism. Publié dans Biophysical J., 2189-Plat, 2001.
17. **LOUSSOUARN G.**, PIKE L.J., ASHCROFT F.M., MAKHINA E.N., NICHOLS C.G. Dynamic sensitivity of K_{ATP} channels to ATP. Publié dans Biophysical J., 27-Plat, 2001.
16. ROSE T., **LOUSSOUARN G.**, PHILLIPS R., NICHOLS C.G. Exploring the pore structure of the inwardly rectifying Kir6.2 channel. Publié dans Biophysical J., 65-Plat, 2001.
15. **LOUSSOUARN G.**, ROSE T., MAKHINA E.N., NICHOLS C.G. Conformational flexibility of inward rectifier K channel pore. Publié dans Biophysical J., 2027-Pos, 2000.
14. **LOUSSOUARN G.**, MAKHINA E.N., ROSE T., NICHOLS C.G. Structure and dynamics of the pore of inward rectifier of K_{ATP} channels. Congrès de l'American Physiological Society. Biology of Potassium Channels: From Molecules to Disease. Publié dans The Physiologist 42, 4, 1999.
13. **LOUSSOUARN G.**, MAKHINA E.N., NICHOLS C.G. Structure of the second transmembrane domain of Kir6.2 revealed by the Substituted Cysteine Accessibility Method. Biophysical J., A73, 1999.
12. MAKHINA E.N., **LOUSSOUARN G.**, NICHOLS C.G. The M2 transmembrane domain of Kir6.2 controls ATP sensitivity of cloned K_{ATP} channels. Publié dans J. Physiol, 509.P, 37P, 1998.
11. **LOUSSOUARN G.**, MAKHINA E.N., NICHOLS C.G. A cysteine residue (C166) in Kir 6.2 is responsible for Cd²⁺ inhibition of cloned K_{ATP} channels. Publié dans J. Physiol, 509.P, 37P, 1998.
10. DEMOLOMBE S., BARO I., BLIEK J., MOHAMMAD PANAH R., **LOUSSOUARN G.**, MANNENS M., WILDE A., BARHANIN J., CHARPENTIER F., PEREON Y., ESCANDE D. The K⁺ channel involved in the long QT Syndrome is a complex made of three different proteins. Publié dans Suppl. to Circulation, 96, I302, 1997.
9. POLLARD H., **LOUSSOUARN G.**, BEHR JP., ESCANDE D. Transportation through the nuclear membrane of cDNA complexed with a novel synthetic vector : polyethylemimine. Pediatric Pulmonology, supp 14, 223, 1997.
8. **LOUSSOUARN G.**, CHARPENTIER C., MOHAMMAD PANAH R., KUNZELMANN K., BARHANIN J., BARO I., ESCANDE D. The chromanol 293B is a blocker of the KvLQT1 current. Publié dans Suppl. to Circulation, 96, I2383, 1997.
6. DEMOLOMBE S., **LOUSSOUARN G.**, LEBLAIS V., FANEN P., ESCANDE D. Fast expression of functional recombinant CFTR protein in human epithelial cells. Publié dans Pediatric Pulmonology, supp 13, 209, 1996.
5. **LOUSSOUARN G.**, BARÓ I., DEMOLOMBE S., MOHAMMAD-PANAH R., ESCANDE D. The cystic fibrosis gene product, CFTR, acts as a potassium channel regulator. Suppl. to Circulation, 94, I473, 1996.
4. **LOUSSOUARN G.**, DEMOLOMBE S., LEBLAIS V., FANEN P., ESCANDE D. Hyperexpression of recombinant CFTR protein altered its sensitivity to cAMP-stimulation. Publié dans Pediatric Pulmonology, supp 13, 215, 1996.
3. TAUC M., BIDET M., **LOUSSOUARN G.**, POUJEOL P. CFTR expression and cAMP sensitive chloride conductance in primary cultures of rabbit renal tubules. XIIIth International Congress of Nephrology, Juillet 1995.

2. MOHAMMAD-PANNAH R., **LOUSSOUARN G.**, ESCANDE D., BARÓ I. CFTR regulates ORCC channels in transfected CFPAC-1 epithelial cells. Publié dans *Pediatric Pulmonology*, supp 12, 181, 1995.
1. **LOUSSOUARN G.**, DEMOLOMBE S., ESCANDE D., BARÓ I. Retroviral CFTR gene transfer to CF epithelial cells influences cAMP-activated K⁺ currents. Publié dans *Pediatric Pulmonology*, supp 12, 181, 1995.