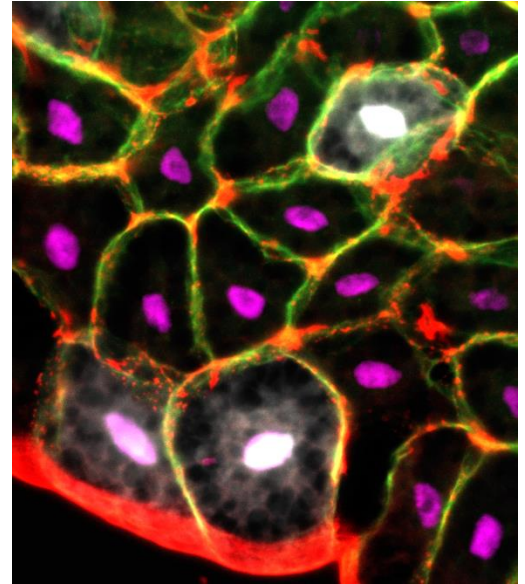


## Offre de poste d'ingénieur d'étude

Patrick Jouandin, équipe Interactions métabolisme-épigénétique chez la *Drosophile*, jeune chef d'équipe ATIP-AVENIR  
CDD 12 mois renouvelable

### Research background & project:

Metabolism is intricately linked with epigenetics. On the one hand, the epigenetic machinery controls chromatin's architecture that governs the expression program of metabolic enzymes. On the other hand, specific metabolic products can act as epigenetic modifiers that can reprogram transcription. This interplay is involved in development, cancer and the immune response, and unravelling this regulatory network will have immediate impact on understanding human health. However, the mechanisms at play and their roles *in vivo* are largely unclear. One challenge has been a lack of genetic models amenable to the systematic characterization of the complex crosstalk between epigenetic and metabolic pathways. Another is that because alteration of the epigenome triggers global transcriptional changes, it is particularly difficult to link a set of specific target genes to a measurable phenotype *in vivo*. To address this, we have established a genetic model in *Drosophila* that allows for medium-throughput screening *in vivo* to systematically interrogate the metabolism-epigenetics interaction during development and inflammation. Our screening results already identified numerous metabolic enzymes and epigenetic factors implicated in these processes, awaiting functional characterization. Our projects fall along three research aims: i) characterize the chromatin landscape bound by epigenetic factors and identify direct metabolic targets, ii) characterize how chromatin remodelers regulate flux through specific metabolic pathways, and iii) identify metabolites secreted from the adipose tissue that regulate development and the inflammatory response. Overall, we aim to decipher, *in vivo*, a complex regulatory network of conserved genes and metabolites that are pathophysiologically relevant in humans.



### Profil :

Recherche d'une personne motivée, sympathique et travailleuse avec un esprit d'équipe et capable de travailler en collaboration. Le sens de l'organisation est un prérequis. Le/la candidat(e) aura pour mission de m'assister dans la mise en place du laboratoire, participer au développement/transfert de technologie ainsi que s'impliquer dans un projet de recherche. Une expérience préalable avec les techniques de bases en biologie moléculaire et/ou en biochimie est requise. Une expérience avec le model *Drosophila* est appréciée mais pas nécessaire.

**Applications:** Envoyer CV et lettre de motivation à [patrick.jouandin@inserm.fr](mailto:patrick.jouandin@inserm.fr).

### Key publications:

1. Gu X\*, Jouandin P\*, Lalgudi PV, Binari R, Valenstein ML, Reid MA, et al. Sestrin mediates detection of and adaptation to low-leucine diets in *Drosophila*. *Nature*. 2022 Aug;608(7921):209–16.
2. Jouandin P, Marelja Z, Shih YH, Parkhitko AA, Dambowsky M, Asara JM, et al. Lysosomal cystine mobilization shapes the response of TORC1 and tissue growth to fasting. *Science*. 2022 Feb 18;375(6582):eabc4203.
3. Parkhitko AA, Jouandin P, Mohr SE, Perrimon N. Methionine metabolism and methyltransferases in the regulation of aging and lifespan extension across species. *Aging Cell*. 2019 Dec;18(6):e13034.
4. Ghiglione C\*, Jouandin P\*, Cérézo D, Noselli S. The *Drosophila* insulin pathway controls Profilin expression and dynamic actin-rich protrusions during collective cell migration. *Development*. 2018 Jul 30;145(14):dev161117.
5. Jouandin P, Ghiglione C, Noselli S. Starvation induces FoxO-dependent mitotic-to-endocycle switch pausing during *Drosophila* oogenesis. *Development*. 2014 Aug;141(15):3013–21.

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Campus Val d'Aurelle

208 rue des Apothicaires  
34298 Montpellier Cedex 5 - FRANCE

E-mail : [patrick.jouandin@inserm.fr](mailto:patrick.jouandin@inserm.fr)

Site Internet : <http://www.ircm.fr>