

# Lignée ORGAPRED N°OV-076\_T

Le MIAOU (**M**inimal **I**nformation **A**bout an **O**rganoid and its **U**se):

Eléments descriptifs permettant à l'homme de l'art de reproduire une expérience de fabrication, de caractérisation et d'étude fonctionnelle d'organoïdes

Le MIAOU sert à identifier les informations présentes (la réponse Oui/Non est la plus importante) et à évaluer la qualité de leur description pour la reproductibilité.

## A) SOURCE MATERIAL MATERIEL SOURCE

|                                                                                                                                               |     |
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| Informed consent obtained<br>Consentement adapté au but de la recherche                                                                       | Yes |
| Collection declaration<br>Déclaration de collection- CODECOH*                                                                                 | Yes |
| Descriptors: gender, age, anatomical region, diagnostic, viral statut<br>Descriptif : genre, âge, région anatomique, diagnostic, statut viral | Yes |
| Clinical data on the patient<br>Tableau clinique du patient                                                                                   | Yes |

\* : gestion de la COnservation D'Éléments du COrps Humain

## Primary cell of patient (and healthy subjects) and tumors Cellule primaire de patient (et sujets sains) et tumeurs

|                                                                                                                                                              |                                                                                                                         |
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| Genetic identity at arrival (example: DNA sequence, snips, digital PCR, STR, CGH array)<br>Identité génétique à réception                                    | Yes, STR on tumor and organoid sample for quality control                                                               |
| Genetic quality control (example : Karyotype, STR, digital PCR)<br>Contrôle de qualité génétique                                                             | Yes, STR on tumor and organoid sample for quality control                                                               |
| Functional quality (example: differentiation test for pluripotency of iPSCs, permeability tests for intestinal epithelial cells...)<br>Qualité fonctionnelle | Yes, organoid establishment                                                                                             |
| Cell identity after X passages<br>identité cellulaire après X passages                                                                                       | STR analysis on tumor DNA and on organoid DNA at passage 12.                                                            |
| Cell type marker (example : marker name, detection method, target value)<br>Marqueur de type cellulaire                                                      | IHC specific markers on tumor fragment: Ki67 / p53 / PAX8 / WT1 / CXCR4 / H3                                            |
| Number of passages at arrival<br>Nombre de passages à la réception                                                                                           | 0, cells directly obtained from patient's tumor                                                                         |
| Number of possible or required passages before genesis of organoids<br>Nombre de passages possibles ou requis avant genèse des organoïdes                    | 0, considered established at passage 3 and above                                                                        |
| Storage conditions<br>Protocole de conservation                                                                                                              | Yes, between 100 and 1000k cells in 2 mL cryovials, resuspended in 1000 µL in freezing solution (10% DMSO, 90% FBS) and |

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|                                                                                                                                                               | frozen gradually decreasing temperature (1°C/min in CoolCell) to -80°C before long-term storage at -196°C.                            |
| Mutations if genetic disease<br>Mutations si maladie génétique                                                                                                | No mutation found by NGS (5 genes panel)                                                                                              |
| Contamination tests (mycoplasma, bacteriological, fungal)<br>Tests contamination                                                                              | Absence of bacterial or fungal contamination                                                                                          |
| Method of tissue dissociation<br>(production of single-cell material or tissue substructures - example: intestinal crypt)<br>Méthode de dissociation du tissu | Yes, 1h enzymatic and mechanical dissociation of tumor fragments with Tumor Dissociation Kit, human (Miltenyi) and 100 µm filtration. |

### Storage conditions of the lines or cells Conditions de conservation des lignées ou cellules

|                                                                                                       |                                                                                                                                                                                                                                                                                                                                                             |
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| Master banks, (description of protocols, drift control)<br>Banques mères                              | Yes, BRC of the BACLESSE Centre, NF S96-900 certified                                                                                                                                                                                                                                                                                                       |
| Daughter banks (description of protocols, drift control)<br>Banques filles....                        | No                                                                                                                                                                                                                                                                                                                                                          |
| Storage: freezing and thawing protocol<br>Conservation : protocole de congélation et de décongélation | Freezing: between 100 and 1000k cells in 2 mL cryovials, resuspended in 1000 µL in Recovery Cell Culture Freezing Medium (Gibco) and frozen gradually decreasing temperature (1°C/min in CoolCell) to -80°C before long-term storage at -150°C.<br>Thawing: Fast thawing at 37°C, transfer in fresh culture media before centrifugation and matrix seeding. |
| Storage modalities<br>Modalités de conservation                                                       | Long-term storage at -150°C.                                                                                                                                                                                                                                                                                                                                |

## B) MANUFACTURING OF THE ORGANIDS FABRICATION DE L'ORGANOIDE

### Culture conditions of cells Conditions de culture des cellules

|                                                                                                                                                                         |                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |
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| Composition of culture media, nature, origin and quantities of supplements used (e.g. glucose, serum, antibiotics, growth factors etc. ....)<br>Composition des milieux | Culture in an enriched medium [Advanced DMEM (Gibco) supplemented with 100 UI/mL of penicillin and streptomycin (Gibco), 1% GlutaMAX (Gibco), 1X B27 (Gibco), 10 mM Nicotinamide (Sigma-Aldrich), 1.25 mM N-Acetyl-L-Cysteine (Sigma-Aldrich), 50 µg/mL Primocin (InvivoGen), 5 µM Y27632 (Interchim), 20 ng/mL FGF-10 (PeproTech), 500 nM A-83-01 (PeproTech), 50 ng/mL EGF (PeproTech), 1 ng/ml FGF-basic (PeproTech), 1 µM SB202190 (PeproTech), 1 µM PGE2 (Sigma- |
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|                                                                           | Aldrich), 10% RSP01-conditioned media (Cultrex HA-R-Spondin1-Fc 293 T, Amsbio) and 50% L-WRN- conditioned media (Cultrex L-WRN, Amsbio)] |
| Nature and treatment of the supports<br>Nature et traitement des supports | Yes. 24-well plate Costar: culture treated<br>6-well plate Nunc: culture treated, Nunclon Delta                                          |
| Seeding conditions<br>Conditions d'ensemencement                          | 10 000 cells per matrix drop (mix of 70% BME2 and 30% culture media v/v)                                                                 |
| Frequency of media changes<br>Fréquence des changements de milieu         | Twice a week                                                                                                                             |
| CO <sub>2</sub> / O <sub>2</sub> Concentration                            | Yes, ambient O <sub>2</sub> and 5% CO <sub>2</sub>                                                                                       |

### Generation of organoids (3D): specificities Génération des organoïdes (3D) : spécificités

#### Matrix culture Culture en matrice

|                                                                                                                               |                                                                                                                                                                                                                                                                  |
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| Nature of the matrix (matrigel, hydrogels, hyaluronic acid, human decellularized matrix etc.)<br>Nature de la matrice         | Cultrex Reduced Growth Factor Basement Membrane Extract, Type 2, Pathclear                                                                                                                                                                                       |
| Matrix concentration<br>Concentration de la matrice                                                                           | 70% matrix in 30% media (v/v)                                                                                                                                                                                                                                    |
| Preparation method (temperature, polymerization time, drop or layer structure, etc.)<br>Modalité de préparation               | 50 µL drop of matrix and 30 minutes polymerization at 37°C                                                                                                                                                                                                       |
| Seeding density per matrix volume unit<br>Densité d'ensemencement                                                             | 10 000 cells per matrix drop                                                                                                                                                                                                                                     |
| Volume and number of drops of matrix per unit area in the culture medium                                                      | 50 µL drop per well in 24-well plate, 10 drops of 50 µL per well in 6-well plate                                                                                                                                                                                 |
| Amount of medium depending on the size of the well<br>Quantité de milieu en fonction de la taille du puits                    | 24-well plate: 500µL<br>6-well plate : 2000µL                                                                                                                                                                                                                    |
| Matrix dissociation method for organoid recovery<br>Méthode de dissociation de la matrice pour la récupération des organoïdes | Mechanical dissociation using cold Advanced DMEM media (Gibco) supplemented with 100 UI/mL of penicillin and streptomycin (Gibco), 1% GlutaMAX (Gibco), 1% BSA (Panreac)                                                                                         |
| Method of dissociation of organoids for their expansion<br>Méthode de dissociation des organoïdes pour leur expansion         | Enzymatic dissociation using TrypLE Express Enzyme (1X) solution, 5-15 minutes incubation at 37°C with gentle agitation every 5 minutes, cold Adv DMEM media (1% BSA, see above) is then add to stop the reaction before centrifugation, numération and seeding. |

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*Culture including multiple cell types* *Culture incluant de multiples types cellulaires*

|                                                                                                 |      |
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| Sequence of co-culturing and adaptation of co-culture media<br>Séquence des mises en co-culture | N.A. |
| Proportion of cell types<br>Proportion des types cellulaires                                    | N.A. |

## C) ORGANOID CHARACTERIZATION CARACTÉRISATION DES ORGANOIDES

The detailed characterization is project dependent, however some standards emerge

|                                                                                                                                                                       |                                                                                 |
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| <i>Morphology</i><br><i>Structure</i>                                                                                                                                 |                                                                                 |
| Appearance, size, shape [circularity, tubularity, regularity of contour (budding)]<br>Aspect, taille, forme                                                           | Round organoids with glandular structures                                       |
| Opacity/réfringency<br>Opacité/réfringence                                                                                                                            | Opaque                                                                          |
| Intra and inter-organoid homogeneity<br>Homogénéité                                                                                                                   | Heterogeneity in shape and size<br>Heterogeneous expression of some IHC markers |
| Expected morphological, architectural and ultrastructural features, organization of cell types (identity, proportions, distribution)<br>Particularités morphologiques | N.A.                                                                            |

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| <i>Molecular Characterisation</i><br><i>Caractérisation moléculaire</i>                                                                 |                                                                         |
| Elements of genomics, transcriptomics, metabolomics, proteomics,<br>Eléments de génomique, transcriptomique, métabolomique, protéomique | NGS<br>Genomic Instability Scar (GIScar)                                |
| Expected specific molecular markers, epigenetic characteristics<br>Marqueurs moléculaires                                               | IHC specific markers on organoids: Ki67 / p53 / PAX8 / WT1 / CXCR4 / H3 |

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| <i>Function</i><br><i>Fonction</i>                                                                                                                                                                      |                                                                                                                                                                                                                         |
| Qualitative and (if possible) quantitative functional characteristic<br>Caractéristique fonctionnelle                                                                                                   | The RECAP test was performed according to the method described in Thorel et al. (2025) (automated scoring of RAD51 foci in PDOs).                                                                                       |
| Response to treatments (pharmacological, chemical, physical, hormonal...) the treatment protocol, and evaluation (quantitative or qualitative) of the response are described<br>Réponse aux traitements | Protocol: when PDO reached the size of 75–150 µm in diameter, they will be collected and resuspended in PDO treatment medium (PDO culture medium lacking primocin, Y-27,632 and N-acetylcysteine) with 2% BME2. 200 PDO |

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|  | <p>per well will be seeded in 100 µL volume in a previously coated (1:1 PDTO treatment medium/BME2) white clear bottom 96-well plates (Greiner). Drug solutions will then be prepared in a 2% BME2/PDTO treatment medium, added to each well and plates will be transferred to a humidified 37°C/5% CO<sub>2</sub> incubator. During the treatment, PDTO will be monitored using IncuCyte S3 ZOOM (Sartorius). One week later, ATP levels will be measured by CellTiter-Glo 3D assay (Promega) and luminescence will be quantified using GloMax Discover Microplate Reader (Promega). The half-maximal inhibitory concentration (IC<sub>50</sub>) and the area under the dose-response curve (AUC) will be computed for each PDTO model.</p> <p><b><u>Evaluation of treatment response for:</u></b><br/> Carboplatin = resistant<br/> Paclitaxel = resistant<br/> Olaparib = sensitive<br/> Niraparib = resistant</p> |
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| <i>Traceability, organoid drift</i><br><i>Traçabilité, dérive des organoïdes</i>                                                                                                                                                     |                                                                                                                                                                                          |
| Traceability of components (batches, suppliers etc.. environments, complements)<br><i>Traçabilité des composants</i>                                                                                                                 | Yes, traceability of every components (batches number, expiration dates ...)                                                                                                             |
| Traceability of conditioned media (drift of cells used for conditioning, control of lines as for those at the origin of the organoid), control of at least one of the growth factors)<br><i>Traçabilité des milieux conditionnés</i> | Traceability of every components (batches number, expiration dates ...) used for conditioned media production.<br><br>No systematic growth factors quantification in conditioned medium. |
| Drift criteria (morphological, structural, functional, molecular....) specific to each organoid. Specify indices if applicable<br><i>Critères de dérive</i>                                                                          | Morphological, response to treatments, IHC markers, STR                                                                                                                                  |
| Robustness criterion (same starting cells, same organoid). Specify indices if applicable<br><i>Critère de robustesse</i>                                                                                                             | No                                                                                                                                                                                       |

### D) USE OF ORGANOIDS UTILISATION DES ORGANOIDES

#### Organoid for basic research *Organoïde en recherche fondamentale*

|  |    |
|--|----|
|  | No |
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## Organoid in preclinical research (pharmacology, toxicology, ...) Organoïde en recherche préclinique (pharmacologie, toxicologie, ...)

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| Functional similarity criterion between the organoid and the mimicked organ (battery of controls to be performed with target values)<br>Similarité fonctionnelle     | Patient clinical data are available to assess concordance with organoid-based tests.<br><br>Partial correlation between tumor and organoids markers. |
| Number of usable passages<br>Applicable for: Preclinical development of a drug candidate (IND file) using organoids<br>Nombre de passages exploitables               | Yes, more than 10                                                                                                                                    |
| Number of usable passages<br>Applicable for: Definition of predictive signatures of responses (companion test)<br>Nombre de passages exploitables                    | Yes, more than 10                                                                                                                                    |
| Number of usable passages<br>Applicable for : Validation of a care protocol (specific patient) on a cohort: choice of a therapy<br>Validation d'un Protocole de soin | To be defined                                                                                                                                        |

## Organoid in clinic (personalized, predictive and regenerative medicine, transplantation) Organoïde en clinique

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| GMP certification, total traceability of the components, qualification of the components for Domain 1: Care protocol (specific patient) (validation of the protocol of use of the organoid for the orientation of the therapeutic choice)<br>- Criterion of similarity between the organoid and the biopsy<br>Certification GMP | No |
| GMP certification, total traceability of components, qualification of components for Domain 2: Use in regenerative medicine (same as cell and tissue therapies)<br>- Functionality criteria, safety (Derivation of biological material and evaluation of the risk of cancer)<br>Certification GMP                               | No |