



# Essai Clinique

Généré le 05 mai 2024 à partir de

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| Titre                   | Extended Access of Momelotinib for Subjects With Primary Myelofibrosis (PMF) or Post-polycythemia Vera or Post-essential Thrombocythemia Myelofibrosis (Post-PV/ET MF)  |
| Protocole ID            | GS-US-352-4365  |
| ClinicalTrials.gov ID   | <a href="https://clinicaltrials.gov/ct2/show/study/NCT03441113">NCT03441113</a>   |
| Type(s) de cancer       | Syndrome myélodysplasique   |
| Phase                   | Phase II  |
| Stade                   | Myélofibrose  |
| Type étude              | Traitement  |
| Médicament              | Momelotinib   |
| Institution             | CENTRE UNIVERSITAIRE DE SANTE MCGILL<br> SITE GLEN<br>1001 boul. Décarie , Montréal, QC, H4A 3J1   |
| Ville                   |   |
| Investigateur principal | Dr Jonathan How   |
| Coordonnateur           | Judit Kokai<br>438-888-1582   |
| Statut                  | Actif en recrutement  |
| But étude               | The primary objective of this study is to provide extended access of momelotinib (MMB) in participants with primary myelofibrosis (PMF) or post-polycythemia vera or post-essential thrombocythemia myelofibrosis (Post-PV/ET MF) enrolled in studies GS-US-352-0101 (NCT01969838), GS-352-1214 (NCT02101268), GS-US-352-1154 (NCT02124746), who are currently receiving treatment with MMB (available as 100 mg, 150 mg and 200 mg tablets) and have not experienced progression of disease. |
| Critères d'éligibilité  | <ul style="list-style-type: none"><li>• Currently enrolled in Studies GS-US-352-0101, GS-US-352-1214, or GS-US-352-1154</li><li>• Able to comprehend and willing to sign the informed consent form</li></ul>  |
| Critères d'exclusion    | <ul style="list-style-type: none"><li>• Known hypersensitivity to MMB, its metabolites, or formulation excipients</li></ul>   |