CPI-818, an Oral Interleukin-2-Inducible T-Cell Kinase Inhibitor

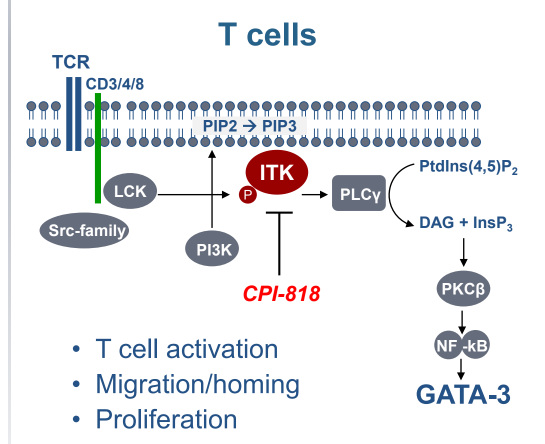
Pre-clinical Characterization and Interim Results of a Phase I/Ib Dose-Escalation Trial in Patients with Relapsed/Refractory T-Cell Lymphoma

Mehrdad Mobasher, MD, MPH Chief Medical Officer



Inhibition of ITK for T Cell Lymphoma

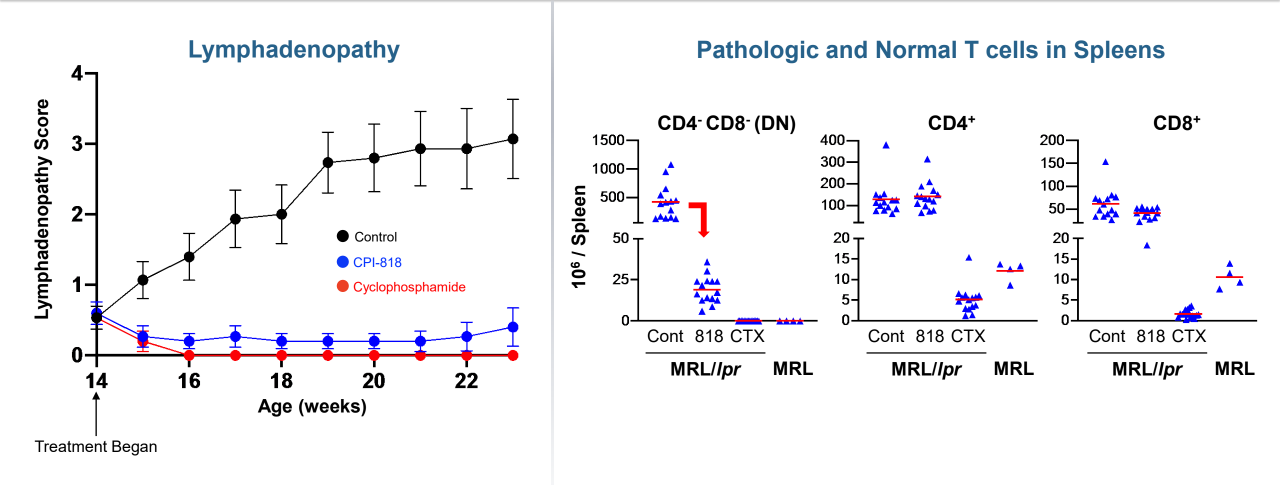
- Rationale for Targeting ITK Signaling in Lymphomas
 - TCR signaling is maintained in most T cell lymphomas
 - Analogous to BCR and B cell lymphomas; ITK is the T cell homologue of BTK and is widely expressed in T cell malignancies
 - Activation of ITK drives NF-κB which drives GATA-3 and survival
 - CTCL and certain PTCLs are thought to be T_H2-driven malignancies
- CPI-818 is a selective, covalent inhibitor of ITK
- Clinical activity observed in canines with CTCL and PTCL



CPI-818 Selectivity Inhibits ITK and Blocks Cellular Signaling CORVUS PHARMACEUTICALS **Kinome Profile and** pPLCγ1 Suppression **Kinases with Cys-442** 000,000,000,00 CPI-818 (µM): 0 TCR αCD3 CPI-818 Kd in nM pPLCγ1 Y783 ITK 2.5 Total PLCγ1 4700 BLK **CPI-818** 9100 BMX ITK **BTK** 1200 **pERK** Suppression 30->10000 **EGFR** MFI Fold Change 25 over unstim) >10000 ERBB2 20. pPLCγ ERBB4 >10000 15-JAK3 2800 10->10000 MKK7 540 TEC **pERK** 0 K_i < 10 nM CP-818 Vehicle CP-818 Vehicle **RLK** 2700 468 Kinases Profiled CD4⁺ T cells CD8⁺ T cells

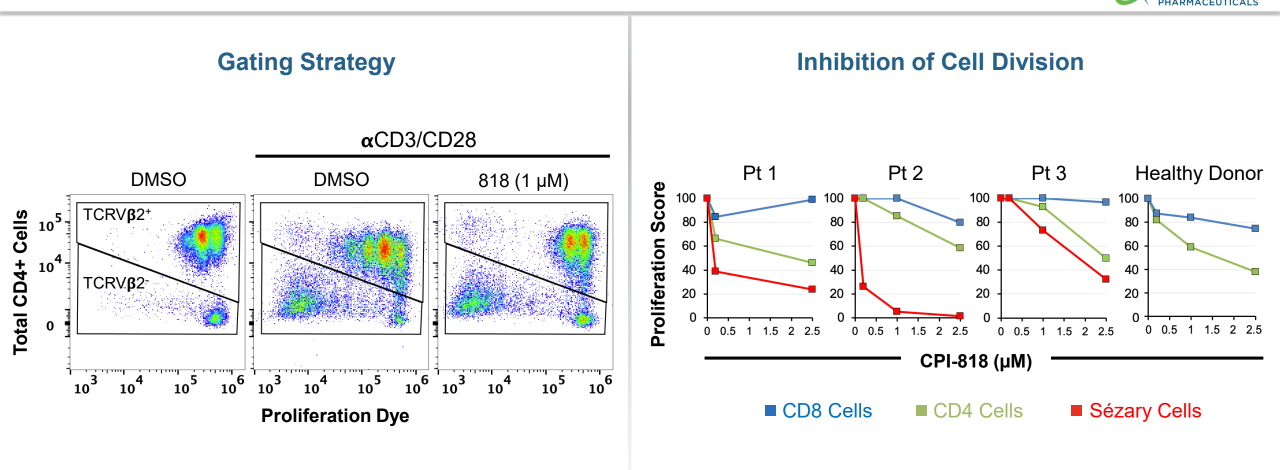
3

CPI-818 Inhibits Lymphoproliferative Disease in Mice



- Fas-/- MRL/lpr mice spontaneously develop lymphoproliferative disease from uncontrolled growth of T cells.
- CPI-818 treatment led to marked regression of lymphadenopathy, with little effect on normal CD4+ and CD8+ T cells.

CPI-818 Preferentially Inhibits Sézary Cells



• Sézary cells were more sensitive than normal CD4+ or CD8+ T cells to the anti-proliferative effect of CPI-818

Spontaneous T Cell Lymphoma in Companion Animals Evaluation of CPI-818 by Prof. Douglas Thamm (CSU)

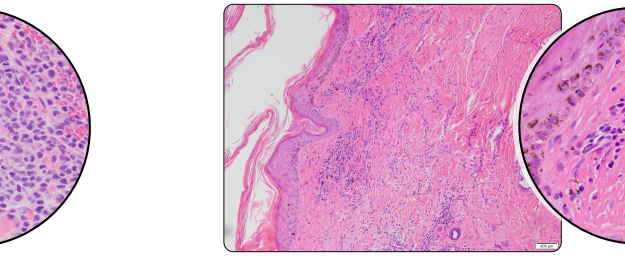


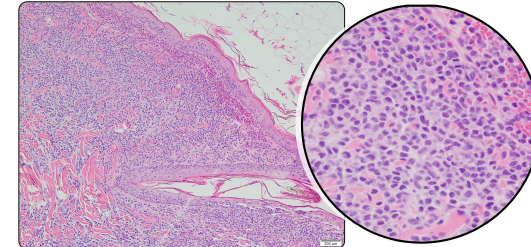


CTCL Patient 11 year old, Male Golden Retriever

4 months

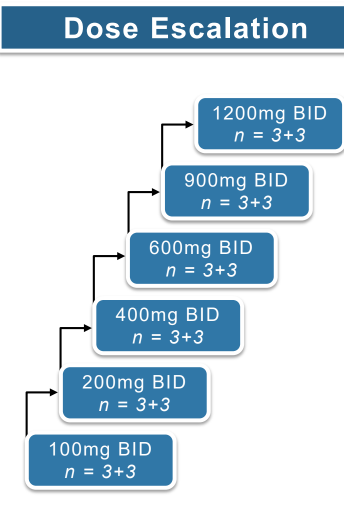






CPI-818-001 Phase 1/1b Clinical Trial Design Dose escalation





Design

- Initial enrollment in dose escalation with 3+3 (+ optional 3) design
- Up to 6 ascending dose levels of CPI-818
- Enroll patients with various types of T-cell lymphoma (PTCL and CTCL) who have progressed on, refractory to, relapsed, or intolerant to at least 2 standard therapies
- Patients will receive CPI-818 orally BID continuously up to sixteen 21-day cycles, until progression or unacceptable toxicity

Objectives

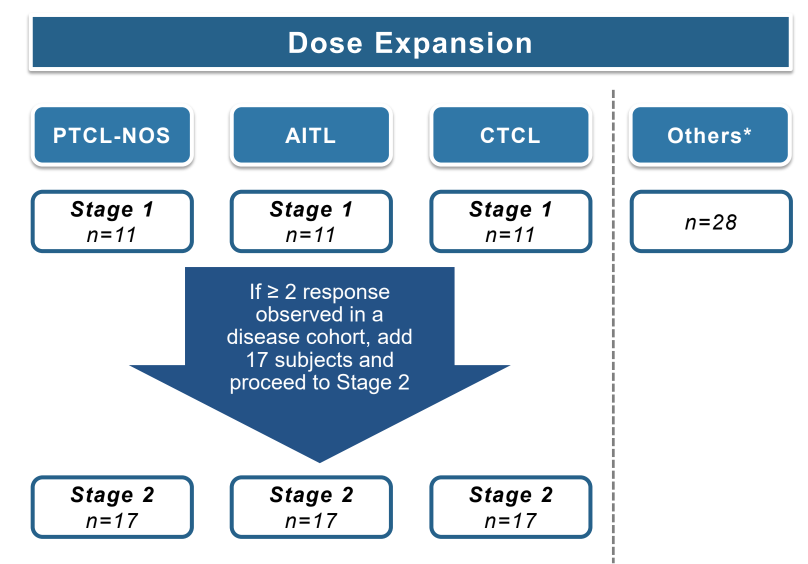
- Primary: To establish Safety/ tolerability and determine MTD or MAD, as well as Expansion Cohort Dose
- Secondary: PK/PD, biomarkers and efficacy

Biomarker Assessments

• ITK in peripheral blood, tissue, cytokines, etc.

CPI-818-001 Trial Dose expansion





* Other types include NKTCL, ALCL, ATLL, etc

CPI-818-001 Study Patient Characteristics



	CPI-818 100mg BID (N=4)	CPI-818 200mg BID (N=3)	CPI-818 400mg BID (N=5)	CPI-818 600mg BID (N=4)
Age (yrs), median (range)	51 (29, 75)	59 (57, 60)	69 (42, 80)	69 (34, 74)
Gender, male N (%)	3 (75%)	1 (25%)	3 (60%)	2 (50%)
No. of prior therapies, median (range)	3 (2, 4)	3 (2, 6)	7 (3, 10)	5 (4, 8)
Histologies	N	N	N	N
Adult T cell leukemia/lymphoma	1	0	0	0
Peripheral T cell lymphoma- NOS	1	1	0	2
Angioimmunoblastic T cell lymphoma	1	1	0	1
Anaplastic large cell lymphoma	1	0	0	0
CTCL (Sézary syndrome)	0	1	4	0
CTCL (Mycosis fungoides)	0	0	1	1

All Adverse Events No grade 3/4 AEs



Adverse Events N (%)	100mg (N=4)	200mg (N=3)	400mg (N=5)	Total (N=12)
Patients with any TEAE	4 (100%)	3 (100%)	3 (60%)	10 (83.3%)
Abdominal distension	1 (25.0%)	0	0	1 (8.3%)
Diarrhea	1 (25.0%)	0	0	1 (8.3%)
Nausea	1 (25.0%)	0	1 (20%)	2 (16.7%)
Retching	1 (25.0%)	0	0	1 (8.3%)
Vomiting	1 (25.0%)	0	0	1 (8.3%)
Chills	1 (25.0%)	0	0	1 (8.3%)
Fatigue	0	1 (33.3%)	0	1 (8.3%)
Pyrexia	0	1 (33.3%)	0	1 (8.3%)
Skin wound (trauma)	0	1 (33.3%)	0	1 (8.3%)
Decreased weight	1 (25.0%)	0	0	1 (8.3%)
Decreased appetite	1 (25.0%)	0	0	1 (8.3%)
Hypercalcaemia	1 (25.0%)	0	0	1 (8.3%)
Hyperuricaemia	1 (25.0%)	0	0	1 (8.3%)
Hypomagnesaemia	1 (25.0%)	0	0	1 (8.3%)
Musculoskeletal pain	0	0	1 (20%)	1 (8.3%)
Headache	0	1 (33.3%)	0	1 (8.3%)
Anxiety	1 (25.0%)	0	0	1 (8.3%)
Cough	0	1 (33.3%)	0	1 (8.3%)
Rash erythematous	0	1 (33.3%)	0	1 (8.3%)
Hyperhidrosis	1 (25.0%)	0	0	1 (8.3%)
Skin Pain	1 (25.0%)	0	0	1 (8.3%)
Pruritus	1 (25.0%)	1 (33.3%)	0	2 (16.7%)
Rash	0	1 (33.3%)	1 (20%)	2 (16.7%)

*Data cut off date: 27Jan2020; 600 mg patients not included as the cohort opened recently

Safety No DLTs



No DLTs observed so far and MTD not reached

• 100mg cohort:

- No treatment related Grade ≥ 2 AEs or SAEs
- Treatment related AEs: diarrhea, nausea, retching (all n=1; Grade 1).
- Other AEs: abdominal pain, vomiting, chills, weight decreased, decreased appetite, hypercalcemia, hyperuricemia, hypomagnesemia, anxiety, hyperhidrosis, pain in skin, pruritus in the setting of disease progression (all n=1)

• 200mg cohort:

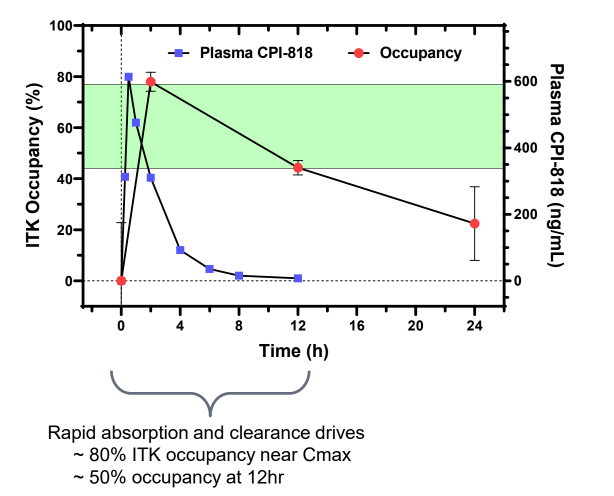
- No treatment related Grade \geq 2 AEs or SAEs
- Treatment related AEs: fatigue, rash (all n=1; Grade 1)
- Other AEs: soft tissue injury, pyrexia, headache, cough, pruritus, rash erythematous due to steroid discontinuation (all n=1)

• 400mg cohort:

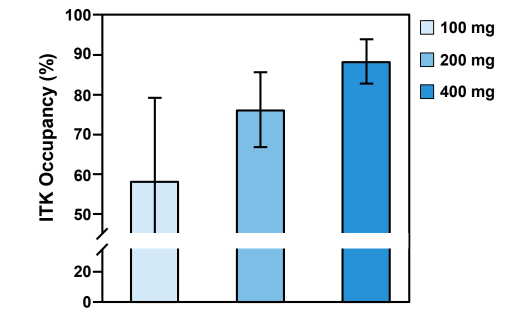
- -4 DLT evaluable pts with no AEs
- Treatment related AEs: nausea (n=1; Grade 1), Grade 2 rash in DLT unevaluable patient (treatment hold
 - 13 days)
- Other AE: Musculoskeletal pain (n=1; Grade 1)

PK and Occupancy Summary from Cohorts 1, 2 and 3 Occupancy is increasing as a function of dose, BID Dosing Required

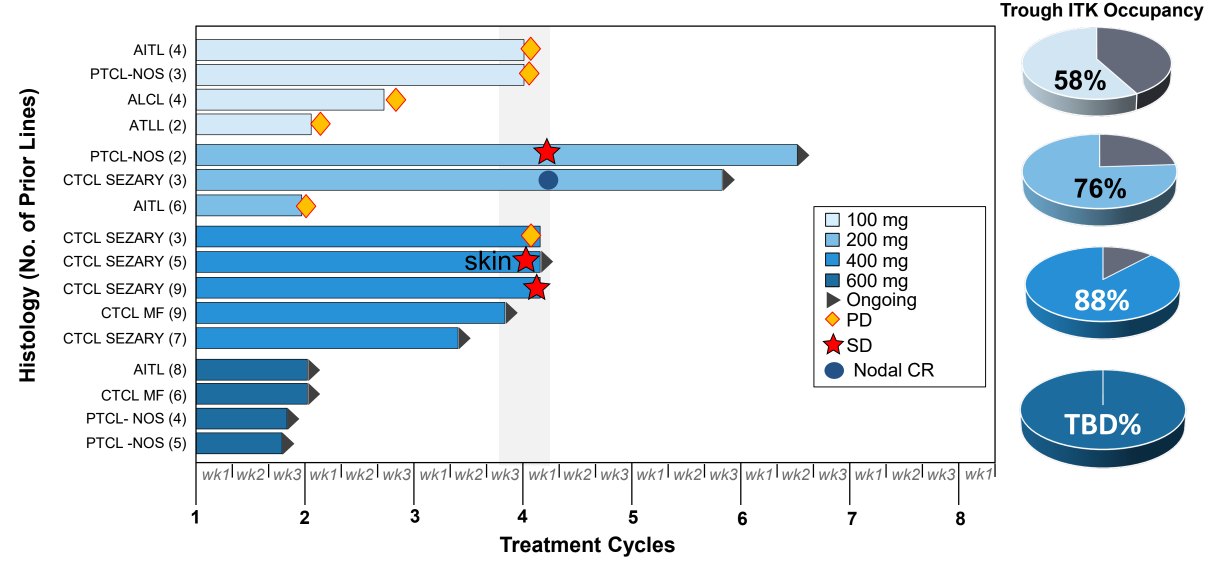
> The Pharmacokinetic profile of CPI-818 and ITK Occupancy in PBMCs following a 100 QD dose



Trough ITK Occupancy for Cycle 1 with BID Dosing of CPI-818



Preliminary Patient Status in Dose Escalation CPI-818-001 Study



median time on treatment to 55 days

CORVUS

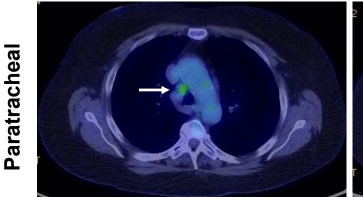
CTCL/SS Patient on 200mg Cohort with Nodal CR Summary



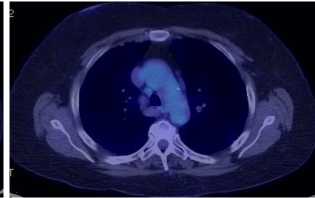
- 60 year old Caucasian female with SS
- FDG avid adenopathy in cervical, axillary, inguinal nodes bilaterally at diagnosis in 2017
- Enrolled in 200mg BID cohort since 21 Oct, 2019
- Screening PET: Small mediastinal (Paratracheal, subcarinal, hilar) and mandibular lymph nodes with moderate FDG avidity
- C4D1 PET Dec 2019 : Interval reduction in the mediastinal nodes with no focus of FDG activity
- Last Visit = C5D1 (13-Jan) remains stable
- mSWAT stable, Sézary cells stable

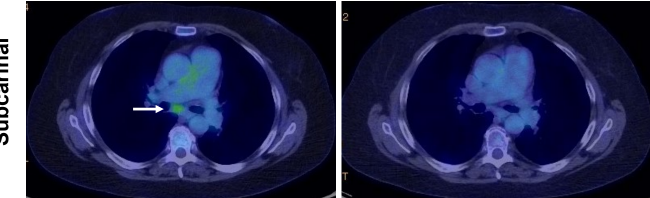
Subcarinal

Pre-Treatment



Post-Treatment(C4D1)





CTCL/SS Patient on 400mg cohort with Skin Improvement Summary



- 80 year old Caucasian male CTCL (Sézary Syndrome)
- Started C1D1 on 400mg BID cohort on 25-Nov-2019
- Screening: no visceral disease or target lesions on CT, absolute Sézary count 5.6x10⁹/L
- Patient reported decreased skin redness with start of treatment after treatment
- C4D1 response assessment: There has been a clinical improvement (mSWAT 92->71)
- Lymphocytosis stable





- CPI-818 is a selective, covalent inhibitor of ITK (sparing RLK and BTK)
- Blocks signal transduction in endpoints downstream of T-cell activation
- Inhibits lymphoproliferative disease in MRL mouse model
- Preferentially inhibits the proliferation of malignant cells sparing normal T cells in blood isolated from sézary patients
- In a companion animal study in dogs with PTCL and CTCL, CPI-818 was well tolerated with evidence for clinical responses
- Interim data from dose escalation part of CPI-818-001 trial shows 100 mg, 200 mg and 400 mg BID doses were well tolerated. Early signs of clinical activity is observed.
- Dose escalation continues

Acknowledgements

