One Mission, Inspired by Patients: Defeat Cancer.™

November 2022





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adversely from those expressed in any forward-looking statements as a result of various factors, including, among others, the success of our corporate strategy and restructuring plan, the success of our commercialization efforts with respect to QINLOCK, including our launch in Germany, our limited experience as a commercial company, our ability to obtain, or any delays in obtaining, required regulatory approvals for our drug and drug candidates, our reliance on sole source thirdparty suppliers, our exposure to criminal sanctions, civil penalties, contractual damages, reputational harm, and diminished profits and future earnings from our relationships with customers and third-party payors that are subject to applicable anti-kickback, fraud and abuse, and other healthcare laws and regulations, and health information privacy and security laws, our failure to obtain or maintain adequate coverage and reimbursement for new or current products, recently enacted and future legislation may increase the difficulty and cost for us to obtain marketing approval of and commercialize our drug candidates, the potential for QINLOCK or any current drug candidates, such as vimseltinib and DCC-3116, or future drug candidates, if successfully developed and approved, to cause undesirable side effects that limit the commercial profile or result in other significant negative consequences for approved products or delay or prevent further development or regulatory approval with respect to drug candidates or new indications, or cause regulatory authorities to require labeling statements, such as boxed warnings, our competition may discover, develop, or commercialize products before or more successfully than we do, our estimates for market opportunities for our approved drug and drug candidates, market acceptance by physicians, patients, third-party payors, and others in the medical community of QINLOCK, and of any future approved drugs, such as vimseltinib or DCC-3116, if approved, our failure to obtain additional marketing approvals in other foreign iurisdictions, the potential for ongoing enforcement of postmarketing requirements for QINLOCK and any drug candidate for which we obtain marketing approval to subject us to substantial penalties, including withdrawal of QINLOCK or any future approved product from the market, if we fail to comply with all regulatory requirements, our ability to complete, and the costs and timing of completing, the development and

commercialization of our drug and drug candidates, our success in enrolling patients in clinical trials, including in our clinical trials of vimseltinib or DCC-3116, the potential for serious adverse events or unacceptable side effects to be identified during the development of our drug or drug candidates, our ability to obtain or, if granted, retain orphan drug exclusivity for our drug or drug candidates; the ongoing effects of the COVID-19 pandemic, our incurrence of significant operating losses since our inception and our expectation that we will incur continued losses for the foreseeable future and may never achieve or maintain profitability, our ability to raise capital when needed, our reliance on third parties to conduct our clinical trials and preclinical studies, our reliance on third parties for the manufacture of our drug candidates for preclinical testing and clinical trials, and for the manufacture of QINLOCK for commercialization and clinical trials, and our ability to enforce our intellectual property rights throughout the world. New risk factors and uncertainties may emerge from time to time, and it is not possible to predict all risk factors and uncertainties. There can be no assurance that the opportunity will meet your investment objectives, that you will receive a return of all or part of such investment. Investment results may vary significantly over any given time period. The appropriateness of a particular investment or strategy will depend on an investor's individual circumstances and objectives. Investors should independently evaluate specific investments. For further information regarding these risks, uncertainties and other factors, you should read the "Risk Factors" section of Deciphera's Quarterly Report on Form 10-Q for the guarter ended September 30, 2022 filed with the Securities and Exchange Commission (the "SEC"), and Deciphera's other SEC filings.

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## ONE MISSION, INSPIRED BY PATIENTS: DEFEAT CANCER™



## SUCCESSFUL COMMERCIAL FRANCHISE AND RESEARCH ENGINE TO FUEL PIPELINE

## QINLOCK®

- Building on successful U.S. launch
- Now approved in nine territories, including the U.S., E.U., and China
- Launch in Germany underway
- Paid access program underway in France

### **Vimseltinib**

- Potential best-in-class product profile
- Estimated \$850M market opportunity in the U.S. for TGCT
- Phase 3 MOTION study enrolling

## **Leader in Autophagy**

- DCC-3116, potential first-in-class ULK inhibitor for cancer in Phase 1
- Significant potential combination opportunity in 70% of cancers

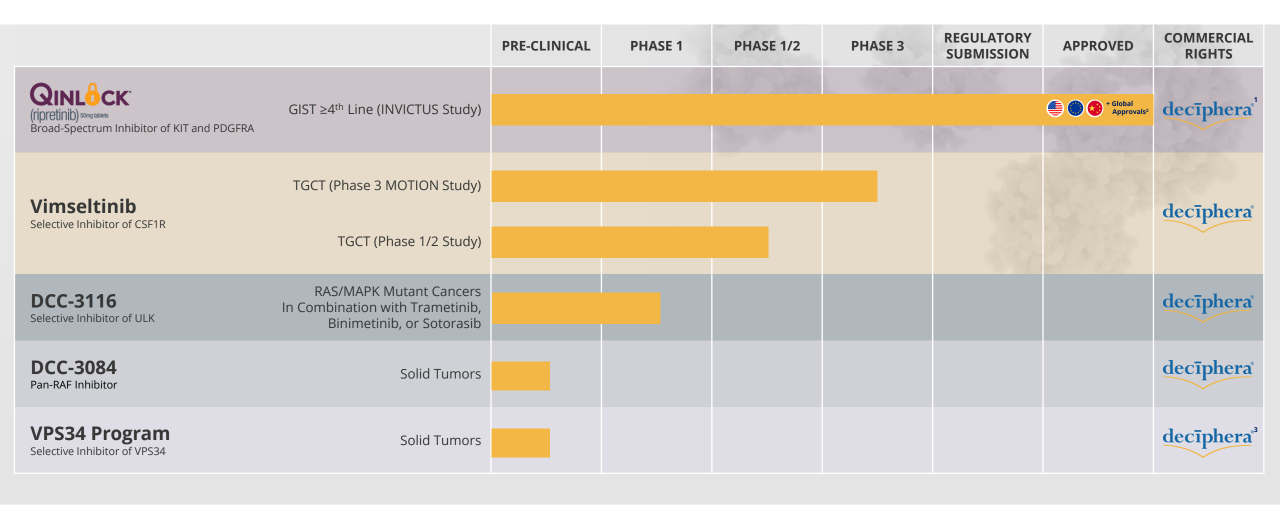
## **Pan-RAF Program**

- Program targeting inhibition of BRAF and CRAF kinases
- Nominated DCC-3084 as our clinical development candidate



Notes: BRAF=proto-oncogene b-RAF; CRAF=proto-oncogene c-RAF; RAF=rapidly accelerated fibrosarcoma; TGCT=tenosynovial giant cell tumor; ULK=unc-51-like autophagy-activating kinase.

## ROBUST PIPELINE OF SWITCH-CONTROL KINASE INHIBITORS





Notes: CSF1R=colony-stimulating factor 1 receptor; GIST=gastrointestinal stromal tumor; KIT=KIT proto-oncogene receptor tyrosine kinase; MAPK=mitogen-activated protein kinase; PDGFRA=platelet-derived growth factor receptor α; RAF=rapidly accelerated fibrosarcoma; RAS=rat sarcoma gene; TGCT=tenosynovial giant cell tumor; ULK=unc-51-like autophagy-activating kinase; (1) Exclusive development and commercialization license with Zai Lab in Greater China for QINLOCK; (2) QINLOCK is approved for 4th line GIST in the United States, Australia, Canada, China, European Union, Hong Kong, Switzerland, Taiwan, and the United Kingdom; (3) Agreement with Sprint Bioscience for exclusive in-license worldwide rights to a research-stage program targeting VPS34.

### STRATEGIC PRIORITIES FOR 2022



QINLOCK® (ripretinib) 50 mg tablets

- Further entrench as standard of care in 4th line GIST in the U.S.
- Successfully launch in key European markets
- Present INTRIGUE data at ASCO Plenary Series Session

#### Vimseltinib

- Rapidly enroll Phase 3 MOTION study
- Present updated clinical data from ongoing Phase 1/2 study

#### **DCC-3116**

- Present initial data from Phase 1 single agent dose escalation study
- Initiate Phase 1 dose escalation study in combination with a MEK inhibitor
- Expand program to include KRAS<sup>G12C</sup> inhibitor combination in NSCLC

## **Proprietary Drug Discovery Platform**

Declare pan-RAF inhibitor development candidate



**Notes:** ASCO=American Society of Clinical Oncology; G12C=single point mutation with a glycine-to-cysteine substitution at codon 12; GIST=gastrointestinal stromal tumor; KRAS=Kirsten rat sarcoma virus; MEK=MAPK/ERK kinase; NSCLC=non-small-cell lung cancer; RAF=rapidly accelerated fibrosarcoma.



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## QINLOCK SUCCESSFUL GLOBAL LAUNCH OF QINLOCK AROUND THE WORLD

## Multiple Global Approvals and Commercial Launches

- Approved in nine territories around the world, including the major markets of the U.S., Europe, and China
- Total revenue of \$36MM in 3Q 2022, which includes QINLOCK product revenue of \$32.3MM and collaboration revenue of \$3.7MM
  - U.S. net product sales of \$24.5MM
  - International net product sales of \$7.8MM
- Direct commercialization in U.S. and E.U.

## QINL CK Total Product Revenue

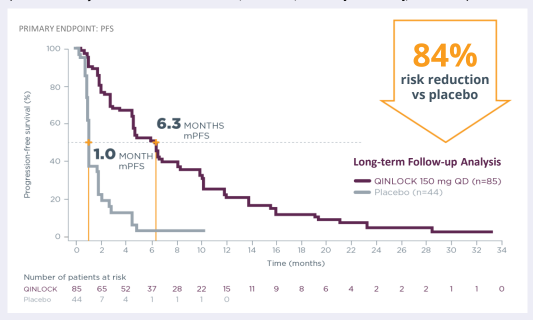




## QINLOCK\* | 4<sup>th</sup> Line Gastrointestinal Stromal Tumor (GIST) CLEAR STANDARD OF CARE FOR THE TREATMENT OF 4<sup>th</sup> LINE GIST

#### **Progression-free survival**

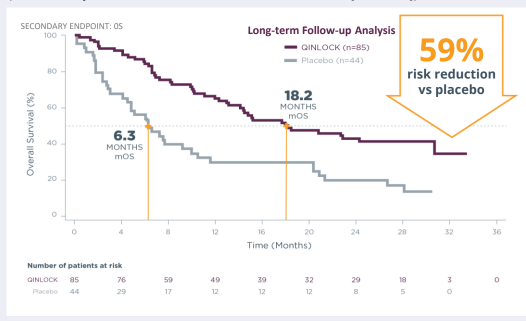
(median PFS of 6.3 months vs. 1.0 month; HR=0.16, 95% CI [0.01-0.27], P<0.0001)



QINLOCK PFS in the primary analysis: 6.3 months vs 1.0 months (HR = 0.15 [95% CI, 0.09 - 0.25], p<0.001)

#### **Overall survival**

(median OS of 18.2 months vs. 6.3 months; HR=0.41, 95% CI [0.26-0.65])



QINLOCK OS in the primary analysis: 15.1 months vs 6.6 months (HR = 0.36 [95% CI, 0.21 - 0.62], p<0.001)

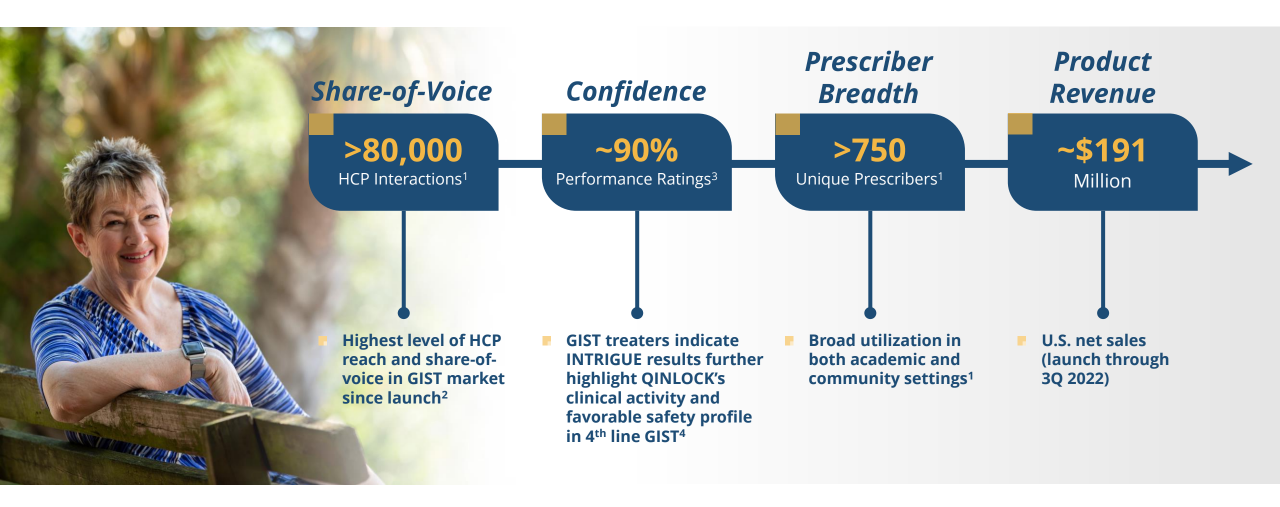
#### **Overall Response Rate:**

- **Long-term follow-up analysis:** 11.8% with QINLOCK vs 0.0% with placebo
- **Primary analysis:** 9.4% with QINLOCK vs 0.0% with placebo (P=0.0504)



**Notes:** Updated data on the INVICTUS Study presented at the ESMO Congress 2021; Full prescribing information is available at <a href="https://www.QINLOCK.com">www.QINLOCK.com</a>; Overall survival data includes all time periods, including dose escalation. Placebo arm includes patients taking placebo who, following progression, were crossed-over to QINLOCK treatment; The long-term follow-up analysis was conducted approximately 19 months from the data cutoff date in the primary analysis and was not powered to show statistical significance; Cl=confidence interval; GIST=gastrointestinal stromal tumor; HR=hazard ratio; mOS=median overall survival; mPFS=median progression-free survival; OS=overall survival; PFS=progression-free survival; QD=daily.

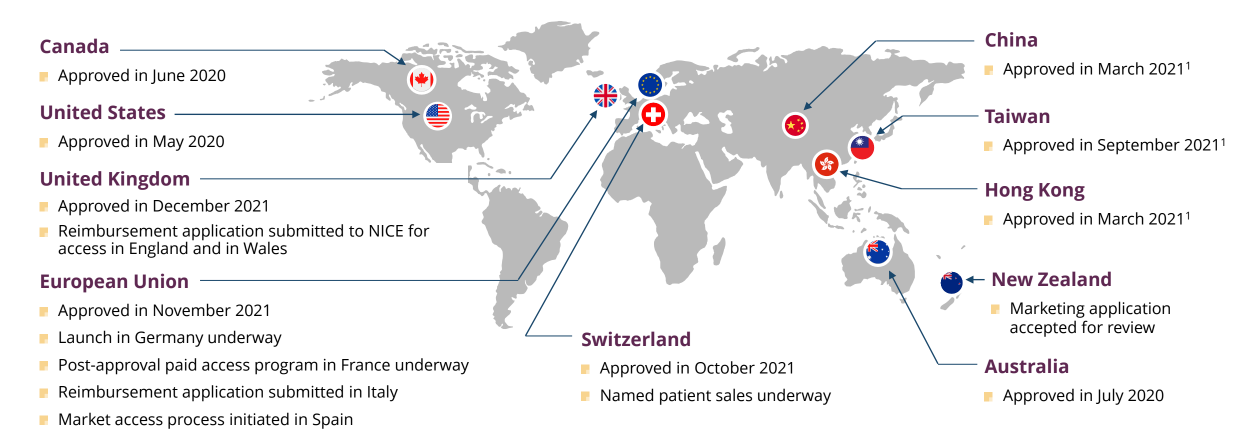
## QINLOCK\* | 4<sup>TH</sup> LINE GASTROINTESTINAL STROMAL TUMOR (GIST) HIGHLY SUCCESSFUL U.S. LAUNCH





**Notes**: GIST=gastrointestinal stromal tumor; HCP=health care provider; QINLOCK launch data represented from May 15, 2020 through September 30, 2022 (1) Internal Deciphera Data (2) Deciphera ATU survey, 2Q 2022 (3) Deciphera ATU survey, 2Q 2022 (3) Deciphera ATU survey, 2Q 2022, ~90% of users rate QINLOCK as performing well to extremely well across key attributes (4) Deciphera market research post-INTRIGUE survey, 4Q 2021.

## Significant progress expanding QINLOCK access to 4th line GIST patients globally





**lotes**: GIST=gastrointestinal stromal tumor; (1) Exclusive development and commercialization license with Zai Lab in Greater China for QINLOCK.

## QINLOCK | 4<sup>TH</sup> LINE GASTROINTESTINAL STROMAL TUMOR (GIST) STRONG LAUNCH IN EUROPE IN 2022 DELIVERING A TOTAL OF \$21.0MM IN INTERNATIONAL NET PRODUCT REVENUE YEAR TO DATE



#### **Fast Patient Access**

- Immediate access and reimbursement in Germany and post approval access program in France underway
- Ongoing discussions with NICE for England and Wales
- Reimbursement submitted in Italy and initiated in Spain
- Infrastructure in place for named patient sales and/or paid access programs as permissible in other European countries



#### Early Revenue Markets

- International QINLOCK net product revenue of \$21.0MM in 2022 driven by EU sales year to date¹
- High demand in Germany and received a "major additional benefit" rating, the first orphan oncology treatment in Germany to receive this rating for its lead indication
- Received unanimous ASMR III rating in France



### **Robust Opportunity**

- Estimated GIST incidence in EU4 +UK comparable to the U.S.: 4,000–6,000 patients
- No other treatment options approved for 4<sup>th</sup> line GIST
- Broad experience with QINLOCK under early access program indicates urgent medical need





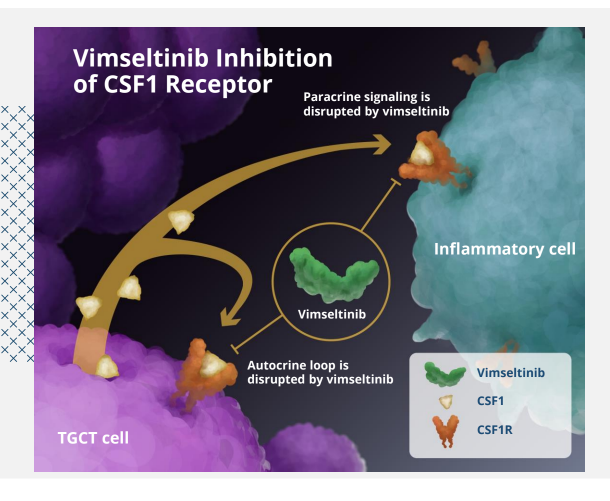
**Notes**: Reimbursement timelines in Europe differ among countries driven by local practices and local requirements; EU5=France, Germany, Italy, Spain, and the United Kingdom; GIST=gastrointestinal stromal tumor. (1) As of September 30, 2022

## VIMSELTINIB



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### ENROLLMENT UNDERWAY FOR PHASE 3 MOTION STUDY IN TGCT



- Vimseltinib is an oral, switch-control TKI specifically designed to selectively and potently inhibit CSF1R
- TGCT caused by translocation in CSF1 gene, coding the ligand for CSF1R
- High unmet medical need in TGCT for effective therapy with improved safety profile
- Positive Phase 1/2 study updates provide strong support for ongoing Phase 3 MOTION study<sup>1</sup>
- Strong strategic fit with GIST based on overlapping KOLs and call-points



Notes: CSF1=colony-stimulating factor 1; CSF1R=colony-stimulating factor 1 receptor; GIST=gastrointestinal stromal tumor; KOL=key opinion leader; ORR=objective response rate; TGCT=tenosynovial giant cell tumor; TKI=tyrosine kinase inhibitor; (1) Data presented at the ESMO Congress 2022.

**Diagnosis** 

Patient burden

**Unmet need** 

## A LOCALLY AGGRESSIVE TUMOR ASSOCIATED WITH SUBSTANTIAL MORBIDITY





## Disease Burden and Unmet Medical Need for TGCT Patients

Due to the slow-growing nature of TGCT and similar symptoms to other conditions like sports injuries and arthritis, patients may have a longer path to diagnosis

In the TOPP registry<sup>1</sup>, patients at baseline commonly reported multiple symptoms, including pain (78%), limited function (64%), swelling (61%), stiffness (55%), and use of analgesics (15%)<sup>2</sup>

- Surgical resection is standard treatment
- High rate of recurrence in diffuse TGCT
- CSF1R inhibition has demonstrated clinical benefit in diffuse TGCT
- Pexidartinib is only approved agent for TGCT patients no longer amenable to surgery (FDA approval in August 2019)
  - FDA label includes boxed warning, REMS, and intensive liver monitoring due to off-target hepatotoxicity risks
  - The EMA adopted the decision of refusal of the Turalio MAA in November 2020
- Unmet medical need for effective CSF1R inhibitor with favorable safety/tolerability profile for TGCT patients



Ankle image courtesy of Tap WD, et al. ASCO 2018, abstract 11502; Knee image courtesy of Tap WD, et al. N Engl J Med. 2015;373:428-437.

Notes: CSF1R=colony-stimulating factor 1 receptor; EMA=European Medicines Agency; FDA=U.S. Food and Drug Administration; MAA=Marketing Authorisation Application; REMS=Risk Evaluation and Mitigation Strategy;
TGCT=tenosynovial giant cell tumor; (1) The TOPP registry was a multinational, multicenter, prospective observational study involving 12 tertiary sarcoma centers in 7 European countries, and 2 US sites; (2) Patients experienced more than or equal to 3 symptoms (52%).

## VIMSELTINIB | ONGOING PHASE 1/2 STUDY OF PATIENTS WITH TGCT STUDY DESIGN

#### PHASE 1 (DOSE ESCALATION)

- A pharmacologically guided 3 + 3 design, to determine the recommended Phase 2 dose (RP2D) and the maximum tolerated dose
- Enrollment in Phase 1 dose escalation is complete

#### COHORT 5 (n=8)

**Loading Dose** 30 mg QD x 5 days

**Dose** 30 mg twice weekly

#### **COHORT 8** (n=12)

**Loading Dose** 30 mg QD x 3 days

**Dose** 10 mg QD

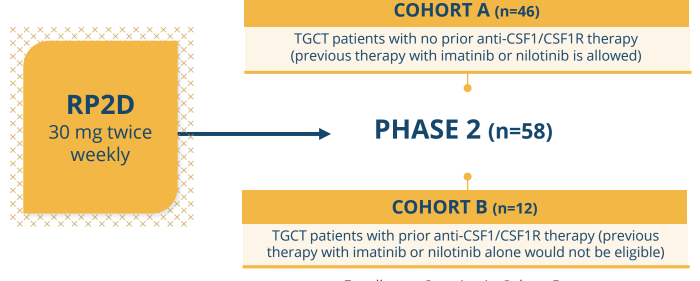
#### **COHORT 9** (n=12)

**Loading Dose** 20 mg QD x 3 days

**Dose** 6 mg QD

#### PHASE 2 (EXPANSION)

- Study to evaluate the safety, tolerability, and preliminary efficacy in two TGCT expansion cohorts
- Recommended Phase 2 dose of 30 mg twice weekly with no loading dose



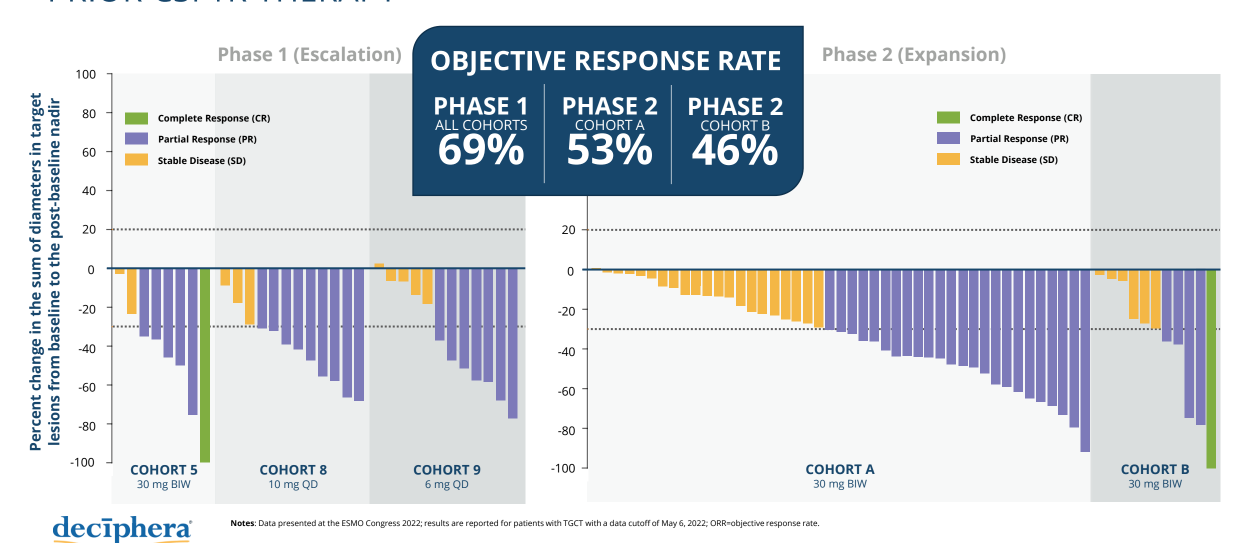




Notes: Data presented at the ESMO Congress 2022; CSF1=colony-stimulating factor 1; CSF1R=colony-stimulating factor 1 receptor; QD=once daily; RP2D=recommended Phase 2 dose; TGCT=tenosynovial giant cell tumor.

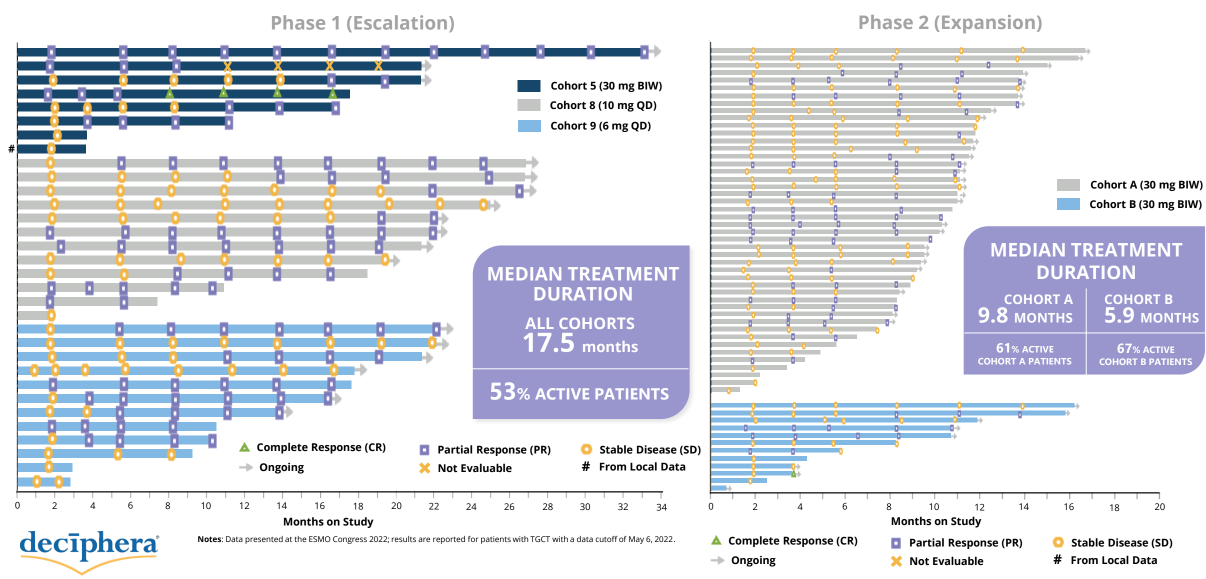
#### VIMSELTINIB | ONGOING PHASE 1/2 STUDY OF PATIENTS WITH TGCT

## ENCOURAGING ANTI-TUMOR ACTIVITY REGARDLESS OF PRIOR CSF1R THERAPY



#### VIMSELTINIB | ONGOING PHASE 1/2 STUDY OF PATIENTS WITH TGCT | PHASE 2 EXPANSION DATA

## INCREASING DURATION OF THERAPY AND DURABLE RESPONSES



#### VIMSELTINIB | ONGOING PHASE 1/2 STUDY OF PATIENTS WITH TGCT

## WELL-TOLERATED IN TGCT PATIENTS

#### **TEAEs in ≥15% of Patients Receiving Vimseltinib**

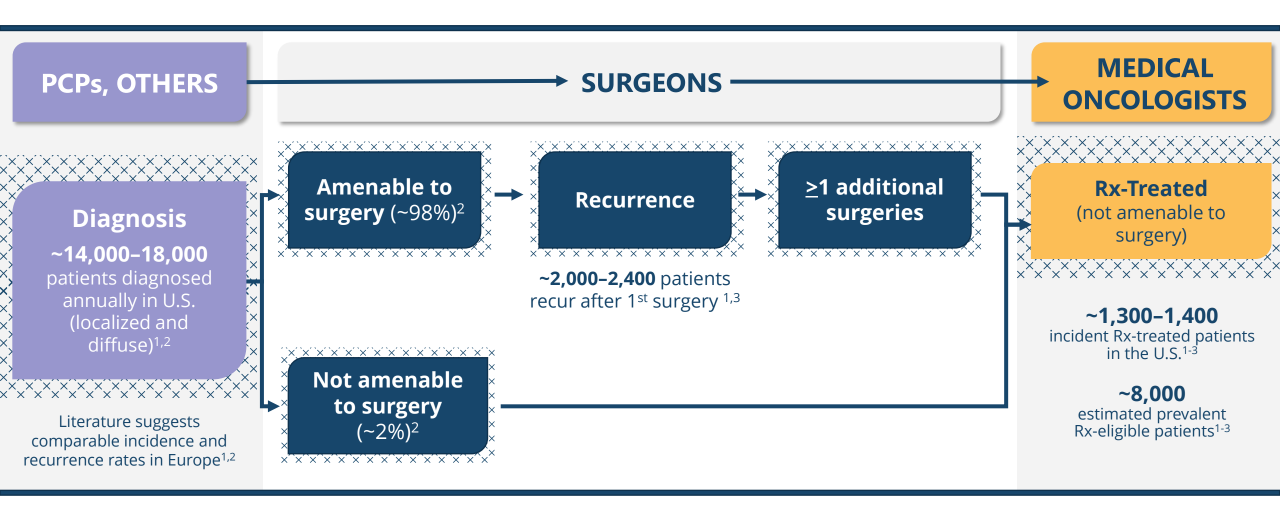
	Pha	se 1	_Pha	ise 2	Phase 1/2 Combined  All Patients (n = 90)		
Preferred term	All Par		All Pa	ntients <sup>1</sup> = 58)			
	All grades	Grade 3/4	All grades	Grade 3/4	All grades	Grade 3/4	
Blood CPK increased	20 (63%)	10 (31%)	34 (59%)	22 (38%)	54 (60%)	32 (36%)	
Periorbital edema	17 (53%)	0	22 (38%)	0	39 (43%)	0	
Fatigue	16 (50%)	0	16 (28%)	0	32 (36%)	0	
AST increased	11 (34%)	4 (13%)	10 (17%)	0	21 (17%)	4 (4%)	
Arthralgia	10 (31%)	1 (3%)	13 (22%)	1 (2%)	23 (26%)	2 (2%)	
Face oedema	9 (28%)	0	10 (17%)	0	19 (21%)	0	
Myalgia	9 (28%)	1 (3%)	15 (26%)	0	24 (27%)	1 (1%)	
Oedema peripheral	9 (28%)	0	9 (16%)	0	18 (20%)	0	
Pruritus	9 (28%)	0	0	0	9 (10%)	0	
Headache	8 (25%)	0	27 (47%)	0	35 (39%)	0	
ALT increased	7 (22%)	1 (3%)	0	0	7 (8%)	1 (1%)	
Diarrhea	7 (22%)	1 (3%)	9 (16%)	0	16 (18%)	1 (1%)	
Lipase increased	7 (22%)	3 (9%)	0	0	7 (7%)	3 (3%)	
Generalized oedema	7 (22%)	0	0	0	7 (7%)	0	
Hypertension	6 (19%)	2 (6%)	0	0	6 (7%)	2 (2%)	
Nausea	6 (19%)	0	19 (33%)	0	25 (28%)	0	
Rash	6 (19%)	0	0	0	6 (7%)	0	
Amylase increased	5 (16%)	2 (6%)	0	0	5 (6%)	2 (2%)	
Constipation	5 (16%)	0	0	0	5 (6%)	0	
Dry skin	5 (16%)	0	0	0	5 (6%)	0	
Paresthesia	5 (16%)	0	0	0	5 (6%)	0	
Rash maculopapular	5 (16%)	0	13 (22%)	1 (2%)	18 (20%)	1 (1%)	
Asthenia	0	0	15 (26%)	1 (2%)	15 (17%)	1 (1%)	

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Notes: Data presented at the ESMO Congress 2022; results are reported for patients with TGCT with a data cutoff of May 6, 2022; Data are presented as n (%) unless otherwise noted; percentages are rounded; ALT=alanine aminotransferase; AST=aspartate aminotransferase; CPK=creatine phosphokinase; TEAE=treatment-emergent adverse event; (1) TEAE cutoff of ≥15% based on all grades for total Phase 1 and Phase 2.

#### VIMSELTINIB | TENOSYNOVIAL GIANT CELL TUMOR (TGCT)

## PATIENT JOURNEY OF TGCT PATIENTS NOT AMENABLE TO SURGERY

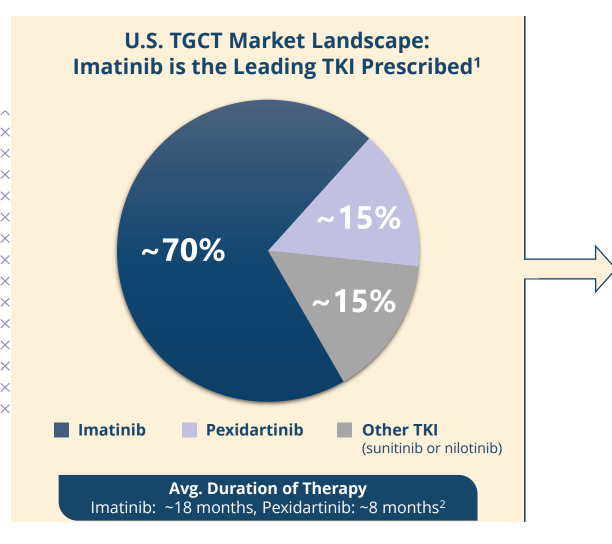




**Notes:** PCP=primary care physician; TGCT=tenosynovial giant cell tumor; (1) Mastboom et al. Acta Orthopaedica. 2017;88(6):688-694; (2) Internal Deciphera estimates based on market research and secondary data; estimates are inherently uncertain; (3) Ehrenstein. J Rheumatol. 2017;44(10):1476-1483.

#### VIMSELTINIB | TENOSYNOVIAL GIANT CELL TUMOR (TGCT)

### TGCT MARKET LANDSCAPE OVERVIEW



## **Existing Product Profiles and Unmet Need**

#### **Imatinib**

- Not FDA or EMA approved for TGCT
- Weak CSF1R inhibitor
- Guideline listed based on retrospective data showing 19% ORR<sup>3,4</sup>

#### **Pexidartinib**

- FDA approved for TGCT, not approved by EMA
- Inhibitor of CSF1R, KIT, FLT3, and PDGFRA/B
- Boxed warning for hepatoxicity, REMS, intensive liver monitoring

### **High Unmet Need**

- Lifelong condition with significant morbidity
- HCPs and patients cite desire for effective therapy without having to sacrifice safety and tolerability<sup>5</sup>

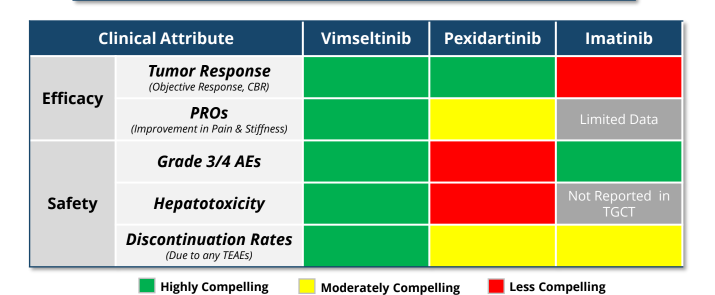


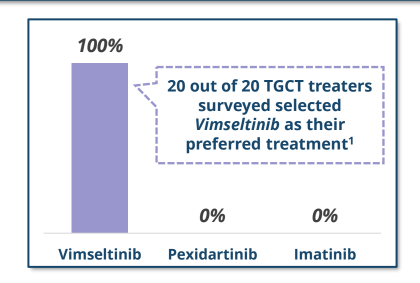
Notes: TGCT=tenosynovial giant cell tumor; TKI = Tyrosine Kinase Inhibitor. (1) Symphony Health IDV Claims; Analysis Period: 7/1/2020 – 6/11/2021; (2) Symphony Health IDV Claims; Analysis Period: 1/1/2008 – 2/1/2021; Patient Qualification: 1/1/2016 – 12/31/2020, Symphony Health IDV Claims; Analysis Period: Q3 2019 – Q32022 and Symphony Health Metys database (estimates calculated from pexidartinib data 2019 – 2022); (3) NCCN Guidelines Version 2.2022 Soft Tissue Sarcoma; (4) Cassier et al Cancer 2012:119:1649-1655; (5) Internal Deciphera market research.

## MARKET RESEARCH HIGHLIGHTS THE POTENTIAL FOR VIMSELTINIB TO DELIVER A BEST-IN-CLASS PROFILE AND ESTABLISH A NEW STANDARD OF CARE IN TGCT

#### **Relative Scoring of Key Product Attributes**

### **Preferred Systemic Treatment For TGCT**





Clinical Profile: "This is very impressive data. Well over half the patients responded to the medication with excellent outcomes. Pain measurements and mobility scores are quite remarkable compared to anything currently available The toxicity profile shows this is a very safe drug as well" – Onc

Efficacy: "It's great to see that the best overall response numbers improve over long-term. In fact, this would probably lead me to keep some patients on it longer than I usually would." – Onc

**Safety:** "Black box warnings are usually at the top of the list of patient concerns. Not having one will be reassuring for them that this is safe to use in the short-term and the long-term." – Onc

**Treatment Choice:** "[Vimseltinib] is clearly superior to the other two products. It has better efficacy and safety data, which is key" – Onc

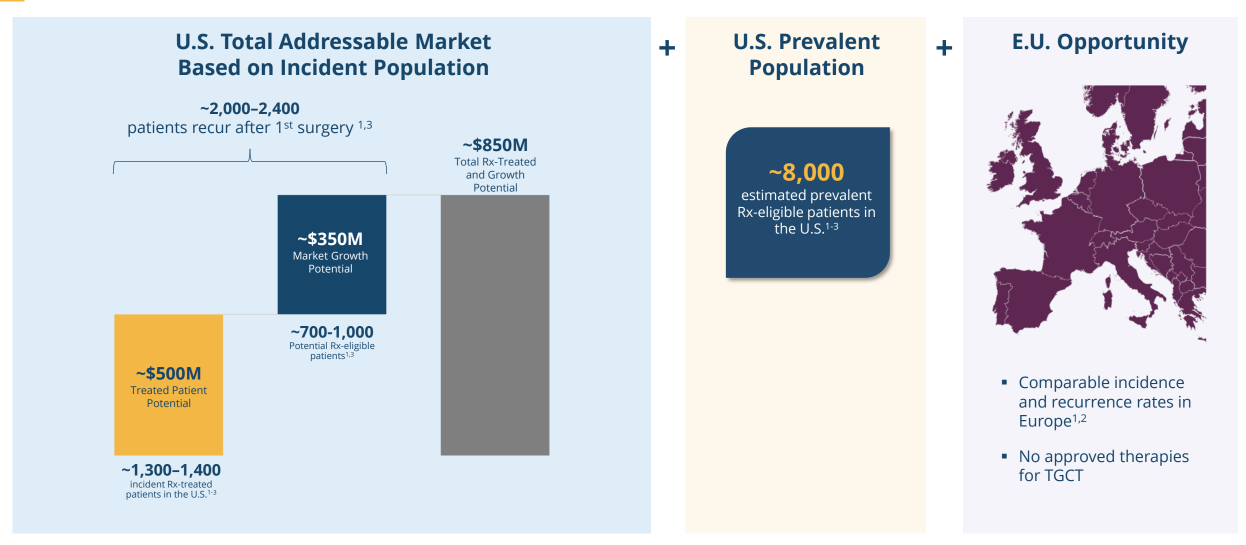
"I would give [vimseltinib] to all my future TGCT patients" - Onc

#### TGCT Treater Sentiments on Vimseltinib Profile



Notes: Qualitative market research conducted by Deciphera in Aug 2022 comparing target product profiles of vimseltinib (blinded), pexidartinib, and imatinib based on USPIs and published/presented literature for each product's efficacy and safety in TGCT diagnosed patients (n=20 HCPs). Hepatotoxicity as defined by increased AST/ALT rates. CBR = Clinical Benefit Rate, PROs = Patient Reported Outcomes, AEs = Adverse Events, TEAEs = Treatment-Emergent Adverse Events. (1) HCP asked to select their preferred (only 1) product to systemically treat TGCT patients based on current clinical profiles.

## SIGNIFICANT OPPORTUNITY TO BENEFIT PATIENTS WITH TGCT GLOBALLY

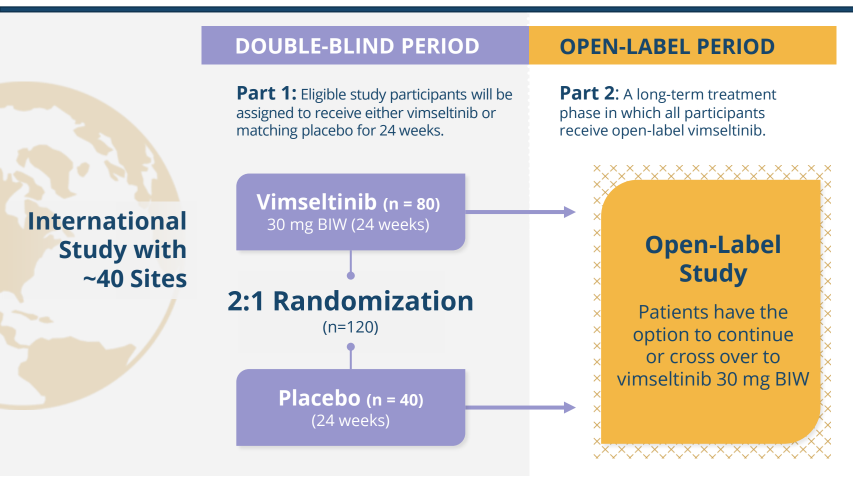




**Notes:** TGCT=tenosynovial giant cell tumor; (1) Mastboom et al. Acta Orthopaedica. 2017;88(6):688-694; (2) Internal Deciphera estimates based on market research and secondary data; estimates are inherently uncertain; (3) Ehrenstein. J Rheumatol. 2017;44(10):1476-1483. Total Addressable Market calculated by estimated patient incidence x 18 months duration x current pexidartinib WAC price and assumes opportunity at steady state.

#### VIMSELTINIB MOTION STUDY | PHASE 3 STUDY OF PATIENTS WITH TGCT

## A MULTICENTER, RANDOMIZED, PLACEBO-CONTROLED, DOUBLE-BLIND STUDY



### **Phase 3 MOTION Study**

will assess the efficacy and safety of vimseltinib for the treatment of patients with TGCT not amenable to surgery

### **Primary Endpoint:**

Objective response rate (ORR) at week 25

### **Key Secondary Endpoints:**

- ORR per tumor volume score
- Range of motion (ROM)
- Patient-reported outcomes



Notes: BIW=twice weekly; TGCT=tenosynovial giant cell tumor.

#### VIMSELTINIB | TENOSYNOVIAL GIANT CELL TUMOR (TGCT)

## QINLOCK AND VIMSELTINIB OFFER THE POTENTIAL TO CREATE A SARCOMA FRANCHISE WITH SUBSTANTIAL OPERATIONAL SYNERGIES

## ~90% of TGCT prescribers are GIST prescribers already targeted by Deciphera<sup>1</sup>



- Ability to commercialize QINLOCK and vimseltinib with a single, sarcoma-focused field team
- Established relationships with sarcoma prescribers
   DCPH sales force ranked highest by GIST treaters among all companies in GIST market<sup>1</sup>
- Existing commercial infrastructure can be leveraged at launch requiring only incremental investment
- Existing strong relationships with KOL and patient advocacy communities

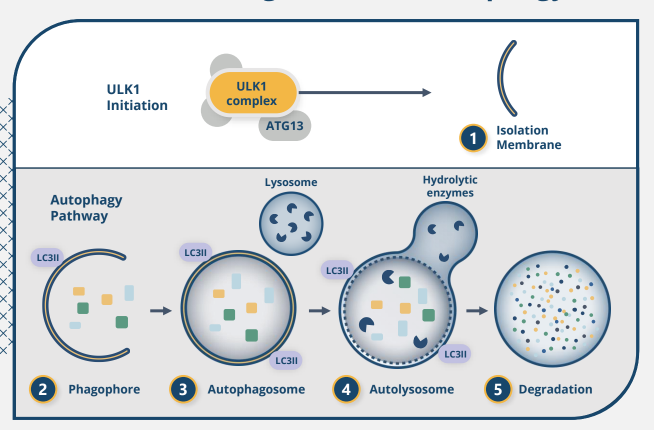


# DCC-3116



### **AUTOPHAGY**: A BROAD RESISTANCE MECHANISM IN CANCER

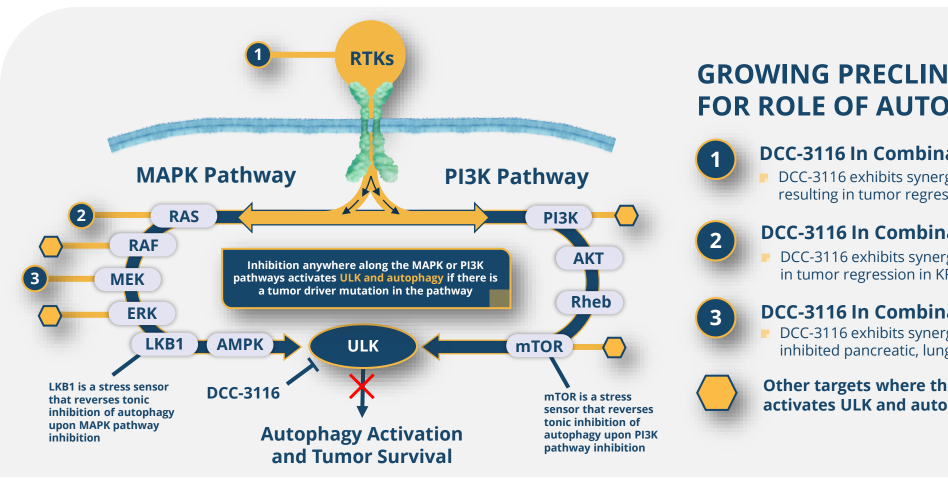
### **ULK: Initiating Factor for Autophagy**



- Autophagy is a catabolic process in which cells recycle components to generate energy
- RTK-, RAS-, and RAF-mutant cancer cells activate autophagy in order to survive stresses or damage **due to anti-cancer therapeutic intervention** with MAPK and PI3K pathway inhibitors
- The **ULK kinase initiates the autophagy pathway** and provides a potential targeted approach to selectively inhibit autophagy in these cancers
- DCC-3116 is a potential first-in-class small molecule designed to inhibit cancer autophagy by inhibiting ULK kinase



## CENTRAL ROLE FOR AUTOPHAGY IN THE RAS/MAPK AND PI3K PATHWAYS



## **GROWING PRECLINICAL VALIDATION** FOR ROLE OF AUTOPHAGY IN CANCER

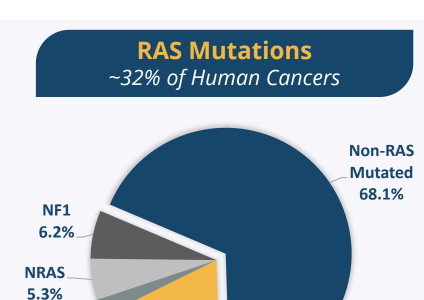
- DCC-3116 In Combination with RTK Inhibition
  - DCC-3116 exhibits synergy with osimertinib and afatinib resulting in tumor regression in EGFR-mutant NSCLC in vivo
- DCC-3116 In Combination with KRAS<sup>G12C</sup> Inhibition
  - DCC-3116 exhibits synergy with sotorasib and adagrasib resulting in tumor regression in KRAS<sup>G12C</sup>-mutant NSCLC in vivo
- DCC-3116 In Combination with MEK Inhibition
  - DCC-3116 exhibits synergy in combination with trametinib and inhibited pancreatic, lung, and melanoma xenograft tumor growth
  - Other targets where therapeutic intervention activates ULK and autophagy



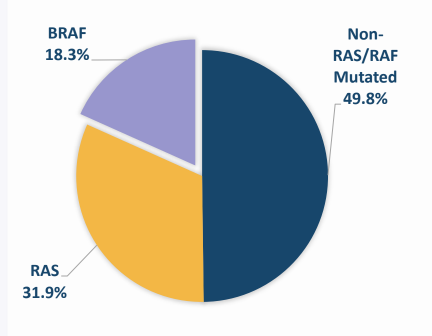
Notes: AKT=protein kinase B; AMPK=AMP-activated protein kinase; EGFR=epidermal growth factor receptor; ERK=extracellular signal-regulated kinase; G12C=single point mutation with a glycine-to-cysteine substitution at codon 12; KRAS=Kirsten rat sarcoma virus; LKB1=liver kinase B1; MAPK=mitogen-activated protein kinase; MEK=mitogen-activated extracellular signal-regulated kinase; mTOR=mammalian target of rapamycin; NSCLC=non-small-cell lung cancer; PI3K=phosphatidylinositol-3 kinase; RAF=rapidly accelerated fibrosarcoma; RAS=rat sarcoma gene; Rheb=ras homolog enriched in brain; RTK=receptor tyrosine kinase; ULK=unc-51-like kinase,

#### DCC-3116

## SIGNIFICANT POTENTIAL OPPORTUNITY EXISTS AS MUTATIONS IN RAS/RAF/RTK SIGNALING OCCURS IN ~70% OF HUMAN CANCERS



## **RAF Mutations** ~18% of Human Cancers





**KRAS** 

18.2%

**HRAS** 

2.2%

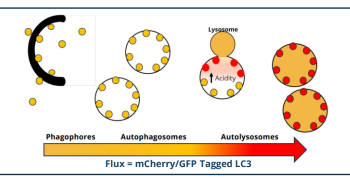
Source: COSMIC (Catalogue of Somatic Mutations in Cancer) Database, v90. **Notes:** RAF=rapidly accelerated fibrosarcoma serine/threonine protein kinase; RAS=rat sarcoma gene; RTK=receptor tyrosine kinase; KRAS=Kirsten rat sarcoma virus; BRAF=proto-oncogene b-RAF; HRAS=Harvey rat sarcoma gene; NRAS=neuroblastoma RAS viral oncogene homolog; EGFR=epidermal growth factor receptor; HER2=human epidermal growth factor receptor 3; PDGFRa=platlet derived growth factor receptor alpha; FLT3=fms-like tyrosine kinase 3; TRK A=Tropomyosin receptor kinase A; TRK B= Tropomyosin receptor kinase B; TRK C=Tropomyosin receptor kinase C; ALK=Anaplastic lymphoma kinase; RET=Rearranged during transfection; FGFR 2=Fibroblast growth factor receptor 2; FGFR 3= Fibroblast growth factor receptor 4; BTK= Bruton tyrosine kinase; cMET=tyrosine-protein kinase Met.

### **RTK Mutations** ~20% of Human Cancers **BRAF** RTK ~20.0% 18.3% Non-RAS/RAF/ **RAS** RTK 31.9% Mutated ~30.0% **RTK Known Tumor Driver Mutations** • TRK A • ALK • FGFR 2 • BCR-ABL • HER2 • PDGFRa • TRK B • ROS • FGFR 3 • BTK • HER3 • FLT3 • TRK C • RET • FGFR 4 cMET exon 14 skipping

#### DCC-3116 | PRECLINICAL DATA

## DCC-3116 OBSERVED PRECLINICALLY TO DURABLY AND POTENTLY INHIBIT AUTOPHAGY INDUCED BY MULTIPLE PATHWAY INHIBITORS

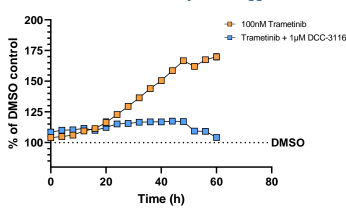
## AUTOPHAGIC FLUX MATURATION



- RTK/RAS/MAPK induces significant autophagy flux
- DCC-3116 can sustainably inhibit this induction

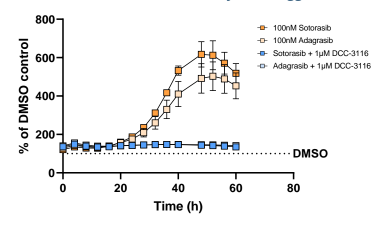
#### DCC-3116 + Trametinib

#### PDAC: Trametinib-induced Autophagy Induction Over Time (mCherry/GFP Tagged LC3)<sup>1</sup>



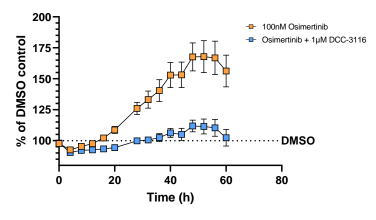
#### DCC-3116 + KRAS<sup>G12C</sup> Inhibitor

## NSCLC: Sotorasib vs. Adagrasib-induced Autophagy Induction Over Time (mCherry/GFP Tagged LC3)<sup>2</sup>



#### DCC-3116 + EGFR Inhibitor

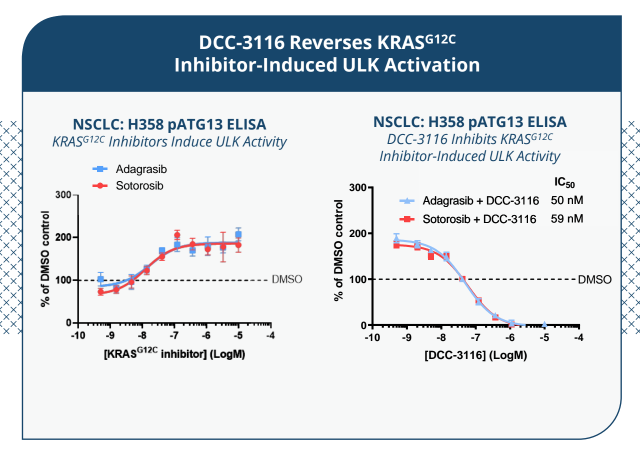
#### NSCLC: Osimertinib-induced Autophagy Induction Over Time (mCherry/GFP Tagged LC3)<sup>1</sup>

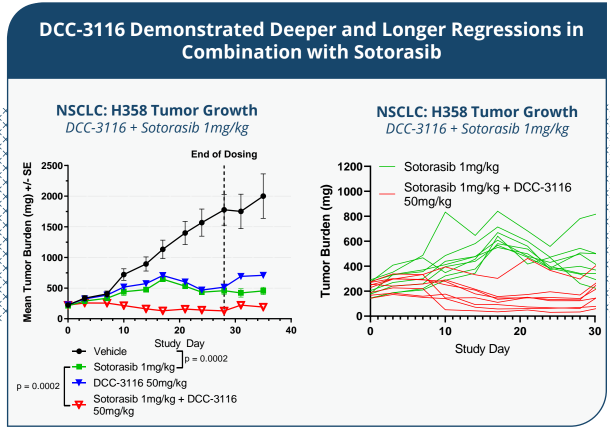




**Notes**: EGFR=epidermal growth factor receptor; G12C=single point mutation with a glycine-to-cysteine substitution at codon 12; KRAS=Kirsten rat sarcoma virus; MAPK=mitogen-activated protein kinase; NSCLC=non-small-cell lung cancer; PDAC=pancreatic ductal adenocarcinoma; RAS=rat sarcoma gene; RTK=receptor tyrosine kinase; (1) data published in Deciphera's Annual Report on Form 10-K for the year ended December 31, 2021; (2) data presented at the AACR Meeting 2022.

## DCC-3116 INHIBITS RAS PATHWAY INHIBITOR-INDUCED ULK ACTIVITY

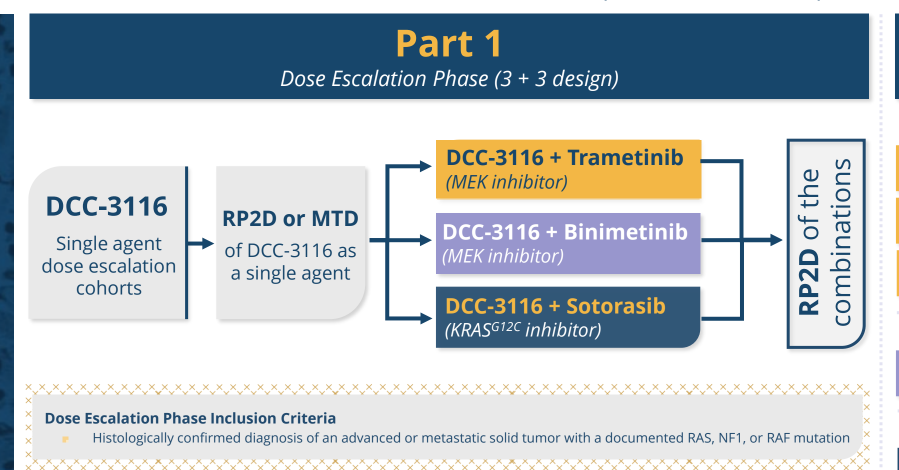






**Notes**: Data presented at AACR 2022; ATG13= Autophagy Related 13; ATG14=Autophagy Related 14: BID=twice daily; MEK=MAPK/ERK kinase; pATG13=phospho-ATG13; pATG14=phospho-ATG14; PBMCs=peripheral blood mononuclear cells; PD=pharmacodynamics; PK=pharmacokinetics; ULK=unc-51-like autophagy-activating kinase.

## MULTICENTER, OPEN-LABEL, FIRST-IN-HUMAN STUDY AS A SINGLE AGENT AND IN COMBINATION TRAMETINIB, BINIMETINIB, OR SOTORASIB



## Part 2

Dose Expansion Phase

DCC-3116 + Trametinib

2<sup>nd</sup> Line PDAC<sup>1</sup>

(KRAS-driven)

3<sup>rd</sup>-5<sup>th</sup> Line NSCLC<sup>2</sup>

(RAF/RAS-driven)

≥3<sup>rd</sup> Line CRC<sup>2</sup>

(RAF/RAS-driven)

DCC-3116 + Binimetinib

2<sup>nd</sup>–3<sup>rd</sup> Line Melanoma<sup>3</sup>
(NRAS-driven)

DCC-3116 + Sotorasib

2<sup>nd</sup>–4<sup>th</sup> Line NSCLC<sup>5</sup> (KRAS<sup>612C</sup>-driven)



Notes: CRC=colorectal cancer; G12C=single point mutation with a glycine-to-cysteine substitution at codon 12; KRAS=Kirsten rat sarcoma virus; MEK=MAPK/ERK kinase; MTD=maximum tolerated dose; NRAS=neuroblastoma RAS viral oncogene homolog; NSCLC=non-small-cell lung cancer; PDAC=pancreatic ductal adenocarcinoma; RAF=rapidly accelerated fibrosarcoma serine/threonine protein kinase; RAS=rat sarcoma gene; RP2D=recommended Phase 2 dose; (1) with a documented mutation in KRAS; (2) with a documented mutation in KRAS; (2) with a documented mutation in KRAS; NFAS, NF1, or BRAF; (3) with a documented mutation in KRAS; (2) with a documented mutation in KRAS; (3) with a documented mutation in KRAS; (4) with a documented mutation i

### SUMMARY OF INITIAL PHASE 1 STUDY RESULTS

- DCC-3116 was well tolerated at doses from 50 to 300 mg BID, which achieved exposure and ULK1/2 inhibition associated with anticancer efficacy with MEK inhibitors in preclinical studies
- Treatment-related TEAEs were mainly Grade 1/2, except for related asymptomatic and reversible Grade 3 ALT increases

DCC-3116 EXPOSURE APPEARED TO INCREASE DOSE PROPORTIONALLY ACROSS 50 – 300 mg BID

NO DLTS OR TREATMENT-RELATED SAES OBSERVED All DOSES ACHIEVED EXPOSURE AND ULK1/2 INHIBITION ASSOCIATED WITH EFFICACY IN PRECLINICAL STUDIES

MONOTHERAPY RESULTS
DEMONSTRATED STABLE
DISEASE AS BEST
OVERALL RESPONSE

NOV '22 UPDATE

MAXIMUM TOLERATED DOSE NOT REACHED 50 mg BID SELECTED AS STARTING DOSE FOR COMBINATION DOSE ESCALATION

FIRST PATIENT TREATED
IN COMBINATION DOSE
ESCALATION



**Notes:** Data presented at the ESMO Congress 2022; data cutoff: June 9, 2022; ALT=alanine aminotransferase; BID=twice daily; DLTs=dose-limiting toxicities; MEK=MAPK/ERK kinase; PD=pharmacodynamics; PK=pharmacokinetics; SAE=serious adverse event; TEAE=treatment-emergent adverse event; ULK=unc-51-like autophagy-activating kinase.

## TEAEs REGARDLESS OF RELATEDNESS (≥15% OF PARTICIPANTS)

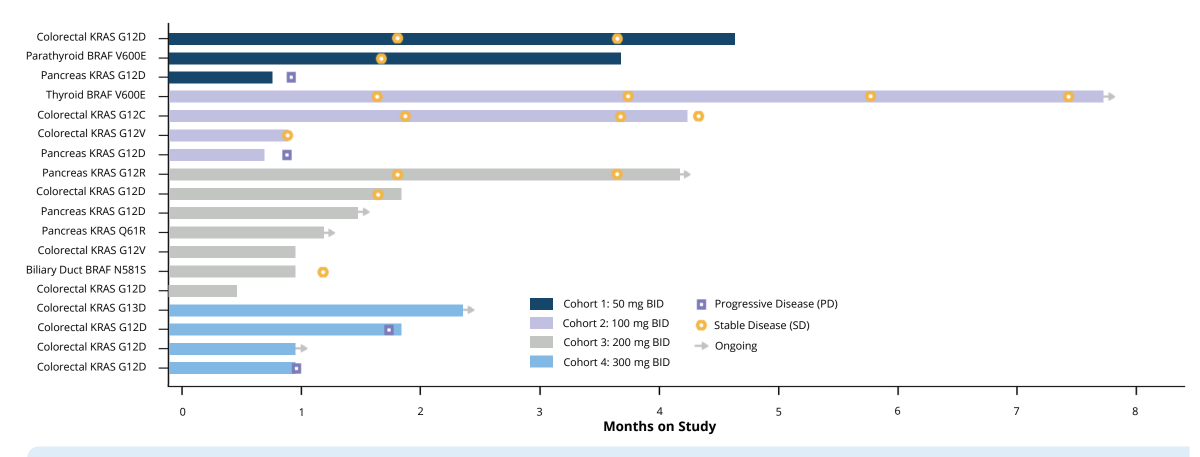
		All Participants							
Preferred term	Cohort 1 50 mg BID (n = 3)		<b>Cohort 2</b> 100 mg BID ( <b>n = 4</b> )		<b>Cohort 3</b> 200 mg BID ( <b>n = 7</b> )		<b>Cohort 4</b> 300 mg BID ( <b>n = 4</b> )		n (%) N = 18
	Grade 1/2	Grade 3	Grade 1/2	Grade 3	Grade 1/2	Grade 3	Grade 1/2	Grade 3	All grades
Fatigue	2	0	1	0	3	0	1	0	7 (39%)
Dehydration	0	0	0	0	2	0	2	0	4 (22%)
ALT increased	0	0	0	0	0	1	1	1	3 (17%)
Anaemia	0	2	0	1	0	0	0	0	3 (17%)
AST increased	0	0	0	0	2	0	1	0	3 (17%)
Decreased appetite	1	0	1	0	0	0	1	0	3 (17%)
Hyponatraemia	1	0	0	0	1	0	1	0	3 (17%)
Nausea	0	0	1	0	0	0	2	0	3 (17%)
Vomiting	1	0	1	0	1	0	0	0	3 (17%)

- No DLTs or treatment-related serious TEAEs were observed
- ► Most treatment-related TEAEs were Grade 1/2; no treatment-related Grade 4 TEAEs
- Two related asymptomatic, reversible Grade 3 ALT increased TEAEs led to dose interruption and reduction



Notes: Data presented at the ESMO Congress 2022; data cutoff: June 9, 2022; ALT=alanine aminotransferase; AST=aspartate aminotransferase; BID=twice daily; DLTs=dose-limiting toxicities; TEAE=treatment-emergent adverse event.

## TREATMENT DURATION AND RECIST v1.1 RESPONSE ASSESSMENTS



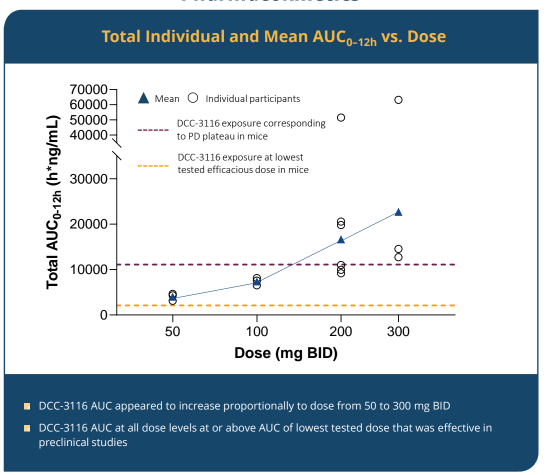
- Best overall response was stable disease
- Disease control rate at week 16 was 29% (4 of 14 response-evaluable participants)



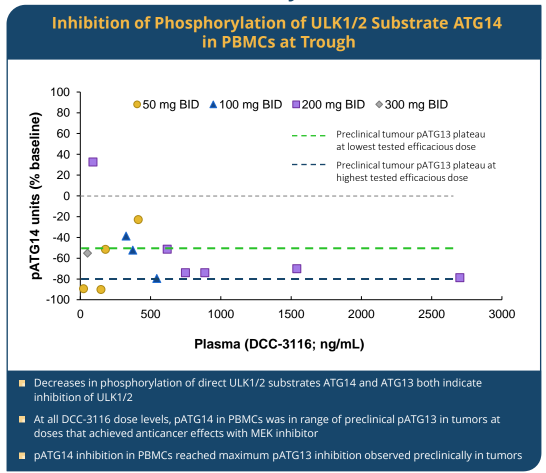
Notes: Data presented at the ESMO Congress 2022; data cutoff: June 9, 2022; BID=twice daily; BRAF=proto-oncogene b-RAF; KRAS=Kirsten rat sarcoma virus.

## INITIAL PK/PD DATA AT ALL DOSE LEVELS IS CONSISTENT WITH PREDICTED EFFICACY BASED ON PRECLINICAL STUDIES

#### **Pharmacokinetics**



### **Pharmacodynamics**

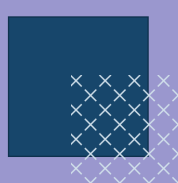




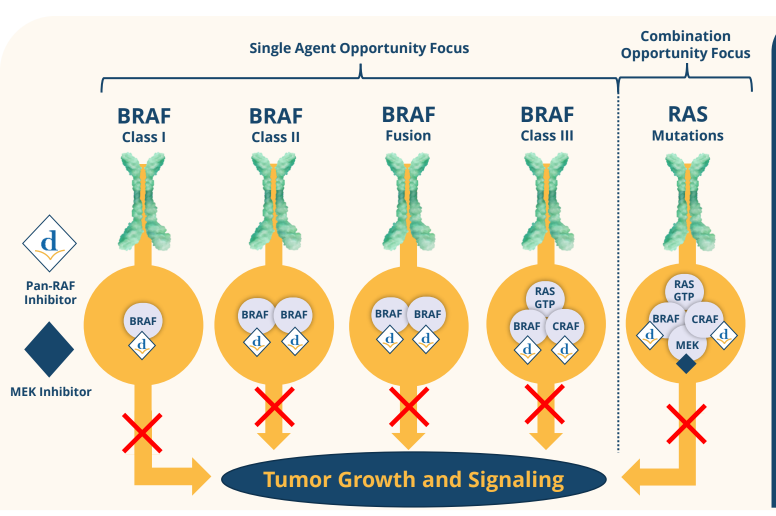
**Notes**: Data presented at the ESMO Congress 2022; data cutoff: June 9, 2022; ATG13= Autophagy Related 13; ATG14=Autophagy Related 14: BID=twice daily; MEK=MAPK/ERK kinase; pATG13=phospho-ATG13; pATG14=phospho-ATG14; PBMCs=peripheral blood mononuclear cells; PD=pharmacodynamics; PK=pharmacodynamics; ULK=unc-51-like autophagy-activating kinase.

# DCC-3084





## POTENTIAL BEST-IN-CLASS PAN-RAF INHIBITOR



- DCC-3084 is a potential best-in-class pan-RAF inhibitor
- Potent and selective inhibitor of BRAF and CRAF kinases targeting Class I, II, and III BRAF mutations as well as BRAF fusions and NRAS mutations
- DCC-3084 designed to be a best-in-class pan-RAF inhibitor based on differentiated pharmaceutical properties and *in vitro/in vivo* profile
- Strong pre-clinical data supports single agent and combination opportunities

Nominated DCC-3084 as the clinical development candidate for the pan-RAF program



**Notes;** BRAF=proto-oncogene b-RAF; CRAF=proto-oncogene c-RAF; GTP=guanosine triphosphate; MAPK=mitogen-activated protein kinase; MEK=mitogen-activated extracellular signal-regulated kinase; RAF=rapidly accelerated fibrosarcoma serine/threonine protein kinase; RAS=rat sarcoma gene.

### EXPECTED MILESTONES ACROSS THE PIPELINE IN 2022





- **✓** Launch QINLOCK in Germany
- **✓** Present INTRIGUE data at ASCO Plenary Series Session
- ✓ Receive authorization for post-approval paid access program in France
- Vimseltinib
  - **✓** Update Phase 1/2 data in TGCT patients
- DCC-3116
  - ✓ Present additional preclinical data
  - ✓ Present Phase 1 single agent dose escalation data
  - ✓ Initiate Phase 1 combination dose escalation cohorts
- Proprietary Drug Discovery Platform
  - **✓** Nominate development candidate for pan-RAF



Notes: ASCO=American Society of Clinical Oncology; RAF=rapidly accelerated fibrosarcoma serine/threonine protein kinase; TGCT= tenosynovial giant cell tumor.

## FINANCIAL HIGHLIGHTS

As of September 30, 2022

Weighted-Average Shares Outstanding<sup>1</sup>

**78.2MM** 

Weighted average shares includes outstanding common stock and common stock issuable upon exercise of prefunded warrants **Cash, Cash Equivalents & Marketable Securities** 

\$372.0MM

Cash Expected to Fund Operating Expenses and CapEx into 2025



## THANK YOU

