

You are cordially invited to attend  
a lunch & learn symposium:

**LONSURF® (trifluridine and tipiracil)  
for the Treatment of Refractory  
Metastatic Colorectal Cancer**



Presented by:

**John Marshall, M.D.**

Georgetown University Medical Center  
Washington, D.C.

Date:

**July 22, 2016**

Location:

**11th Annual New Orleans Summer Cancer Meeting  
Roosevelt Hotel  
Chamber Room  
130 Roosevelt Way  
New Orleans, LA 70112**

Time:

**12:40 PM – 1:45 PM**

**PLEASE VISIT US AT:  
BOOTH #511**

If you have any questions about this program, please contact Jeff Higgins with Taiho Oncology at [jhiggins@taihooncology.com](mailto:jhiggins@taihooncology.com) or Julia Purdue with S Phase at [jpurdue@sphase.com](mailto:jpurdue@sphase.com) or 770-984-5180.

Pursuant to the PhRMA Code on Interactions with Healthcare Professionals, as well as the policies of Taiho Oncology, Inc., attendance at this promotional program is restricted to healthcare professionals (HCPs) within the targeted oncology specialty. Accordingly, spouses and guests are not permitted to attend this program unless they are an HCP within the targeted oncology specialty.

Taiho will report information related to the event, such as your name and the value and purpose of any educational item, meal or other items of value you receive, to the extent required by federal and state laws, as applicable. Please let us know if you are licensed in any state or other jurisdiction, or are an employee or contractor of any organization or government entity that limits or prohibits meals from pharmaceutical companies. HCPs may attend the program and decline a meal. Please note this with your registration and designate this at the venue as you sign-in for the program.

*Please see Indication and Important Safety Information  
on the back and the accompanying full Prescribing Information.*

## INDICATION

LONSURF is indicated for the treatment of patients with metastatic colorectal cancer who have been previously treated with fluoropyrimidine-, oxaliplatin- and irinotecan-based chemotherapy, an anti-VEGF biological therapy, and if RAS wild type, an anti-EGFR therapy.

## IMPORTANT SAFETY INFORMATION

### WARNINGS AND PRECAUTIONS

**Severe Myelosuppression:** In Study 1, LONSURF caused severe and life-threatening myelosuppression (Grade 3-4) consisting of anemia (18%), neutropenia (38%), thrombocytopenia (5%), and febrile neutropenia (3.8%). One patient (0.2%) died due to neutropenic infection. In Study 1, 9.4% of LONSURF-treated patients received granulocyte-colony stimulating factors.

Obtain complete blood counts prior to and on day 15 of each cycle of LONSURF and more frequently as clinically indicated. Withhold LONSURF for febrile neutropenia, Grade 4 neutropenia, or platelets less than 50,000/mm<sup>3</sup>. Upon recovery, resume LONSURF at a reduced dose.

**Embryo-Fetal Toxicity:** LONSURF can cause fetal harm when administered to a pregnant woman. Advise pregnant women of the potential risk to the fetus. Advise females of reproductive potential to use effective contraception during treatment with LONSURF.

### USE IN SPECIFIC POPULATIONS

**Lactation:** It is not known whether LONSURF or its metabolites are present in human milk. There are no data to assess the effects of LONSURF or its metabolites on the breast-fed infant or the effects on milk production. Because of the potential for serious adverse reactions in breast-fed infants, advise women not to breast-feed during treatment with LONSURF and for 1 day following the final dose.

**Male Contraception:** Advise males with female partners of reproductive potential to use condoms during treatment with LONSURF and for at least 3 months after the final dose.

**Geriatric Use:** Grade 3 or 4 neutropenia and thrombocytopenia and Grade 3 anemia occurred more commonly in patients 65 years or older who received LONSURF.

**Renal Impairment:** Patients with moderate renal impairment may require dose modifications for increased toxicity. No patients with severe renal impairment were enrolled in Study 1.

**Hepatic Impairment:** Patients with moderate or severe hepatic impairment were not enrolled in Study 1.

### ADVERSE REACTIONS

**Most Common Adverse Drug Reactions in Patients Treated With LONSURF (≥5%):** The most common adverse drug reactions in LONSURF-treated patients vs placebo-treated patients with refractory mCRC, respectively, were asthenia/fatigue (52% vs 35%), nausea (48% vs 24%), decreased appetite (39% vs 29%), diarrhea (32% vs 12%), vomiting (28% vs 14%), abdominal pain (21% vs 18%), pyrexia (19% vs 14%), stomatitis (8% vs 6%), dysgeusia (7% vs 2%), and alopecia (7% vs 1%).

**Additional Important Adverse Drug Reactions:** The following occurred more frequently in LONSURF-treated patients compared to placebo: infections (27% vs 15%) and pulmonary emboli (2% vs 0%).

Interstitial lung disease (0.2%), including fatalities, has been reported in clinical studies and clinical practice settings in Asia.

**Laboratory Test Abnormalities in Patients Treated With LONSURF:** Laboratory test abnormalities in LONSURF-treated patients vs placebo-treated patients with refractory mCRC, respectively, were anemia (77% vs 33%), neutropenia (67% vs 1%), and thrombocytopenia (42% vs 8%).

**Please see accompanying full Prescribing Information.**