

A Phase 2 Multicenter, Open-Label Study Evaluating the Efficacy and Safety of Dabogratinib (TYRA-300) in Participants With FGFR3-Altered Low-Grade, Intermediate Risk Non-Muscle Invasive Bladder Cancer (SURF302)

Gautam Jayram,¹ Neal D. Shore,² Amirali Salmasi,³ Leslie Aguilar,⁴ Timothy Burn,⁴ Christine Lihou,⁴ Viraj Degaonkar,⁴ Erik Goluboff,⁴ Dinesh Ganapathy,⁴ Seth P. Lerner⁵



BACKGROUND

- Urothelial carcinoma of the bladder is among the 10 most common malignancies worldwide¹; ~75% of cases are diagnosed as NMIBC, a disease with one of the highest patient burdens and one of the most expensive to treat²
- Although NMIBC generally has a favorable prognosis, it is associated with high recurrence rates and a substantial long-term clinical burden²
- IR NMIBC frequently recurs despite available intravesical therapies, highlighting the need for novel targeted treatment strategies³
- Activating *FGFR3* gene alterations are highly prevalent in IR NMIBC, occurring in up to 80% of cases, and represent key oncogenic drivers of urothelial tumorigenesis and recurrence^{1,4}
- Currently available FGFR inhibitors lack FGFR3 isoform specificity and are associated with non-FGFR3 target toxicities resulting from inhibition of FGFR1/2/4, including nail disorders, stomatitis, hyperphosphatemia, central serous retinopathy, and other class-related adverse events^{5,6}
 - Dabogratinib has been designed to be an oral and highly selective inhibitor of activating FGFR3, while minimizing off-target inhibition of FGFR1/2/4 (Table 1)⁷
 - Early clinical data demonstrate a favorable safety profile of dabogratinib at ≤60 mg, with predominantly low-grade TRAEs, no grade ≥3 events, no hyperphosphatemia, and no dose reductions or discontinuations⁸
 - Dabogratinib is being evaluated in patients with *FGFR3*-altered low-grade IR NMIBC in the phase 2 SURF302 study (NCT06995677)⁹

Table 1. In Vitro FGFR Isoform Selectivity of Dabogratinib IC₅₀ (nM)

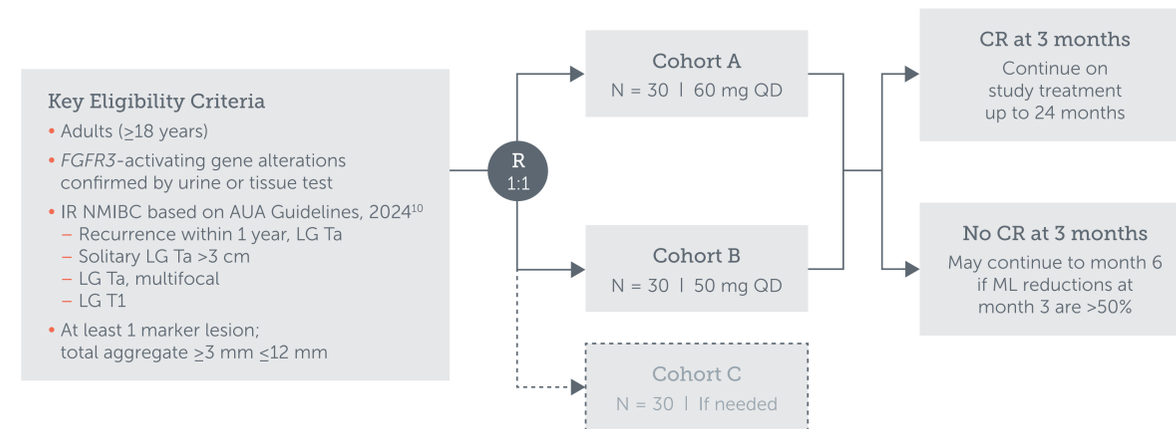
	FGFR3	FGFR1	FGFR2	FGFR4	FGFR1/FGFR3
Dabogratinib	11	278	157	405	25
Erdafitinib	2.6	5.2	3.8	11	2

In engineered Ba/F3 cell assays, dabogratinib shows potent FGFR3 inhibition, with 25-fold selectivity over FGFR1 compared with 2-fold selectivity for erdafitinib. Half-maximal inhibitory concentration values are shown; lower values reflect stronger inhibition.

METHODS

- SURF302 (NCT06995677) is a phase 2, multicenter, open-label, randomized study to evaluate the safety and efficacy of oral dabogratinib QD in adults with *FGFR3*-altered, low grade IR NMIBC⁹
- Participants are assigned to one of 2 parallel dose cohorts:
 - Dose cohort A: dabogratinib 60 mg QD
 - Dose cohort B: dabogratinib 50 mg QD
- An optional third dose cohort may be opened if warranted based on emerging safety and/or efficacy data from cohorts A and B (Figure 1)

Figure 1. Study Design of SURF302



Schematic representation of the study design, illustrating patient randomization, study cohorts, and dabogratinib dosing assignments.

Participant Criteria⁹

Key Inclusion Criteria	Key Exclusion Criteria
<ul style="list-style-type: none"> • Adults ≥18 years of age • Histologically confirmed NMIBC, Ta low grade or T1 low grade • IR NMIBC, defined by AUA criteria 2024¹⁰ (eg, recurrence within 1 year, LG Ta, solitary LG Ta >3 cm, LG Ta multifocal disease, LG T1) • ECOG PS 0-1 • Presence of residual visible marker lesion(s) following diagnostic biopsy/TURBT • Documented activating <i>FGFR3</i> mutation or fusion • No evidence of upper urinary tract or prostatic urethral disease • No prior BCG within 1 year and no intravesical chemotherapy within 8 weeks • Adequate bone marrow, hepatic, and renal function • Ability to swallow oral medication 	<ul style="list-style-type: none"> • Serum phosphate > ULN at screening • Ocular conditions associated with increased risk of eye toxicity • Current evidence of central serous retinopathy or retinal pigmented epithelial detachment of any grade • History of uncontrolled cardiovascular disease • Current or prior muscle-invasive, lymph-node-positive, or metastatic bladder cancer • Tumor involvement of the upper tract or prostatic urethra • Prior treatment with an FGFRi • GI disorder that may affect administration/absorption of dabogratinib • Pregnant or breastfeeding females, or participants planning conception during study participation

Study Endpoints⁹

Primary Endpoint	Secondary Endpoints	Exploratory Endpoints
<ul style="list-style-type: none"> • Complete response rate at 3 months 	<ul style="list-style-type: none"> • Duration of response • Time to recurrence • Recurrence-free survival at 12 and 24 months • Progression-free survival • Incidence and severity of adverse events 	<ul style="list-style-type: none"> • Biomarker analyses, including molecular correlates of response • Longitudinal disease monitoring using clinical and molecular assessment

SURF302 SITE LIST



Site list reflects participating centers as of February 3, 2026; the study is planned for approximately 51 centers across North America, Australia, and Europe.

Abbreviations

AUA, American Urological Association; BCG, bacillus Calmette-Guérin; CR, complete response; ECOG PS, Eastern Cooperative Oncology Group performance status; FGFR, fibroblast growth factor receptor; FGFRi, fibroblast growth factor receptor inhibitor; GI, gastrointestinal; IC₅₀, inhibitory half-maximal concentration; IR, intermediate-risk; ML, marker lesion(s); NMIBC, non-muscle-invasive bladder cancer; QD, once daily; R, randomization; TRAE, treatment-related adverse event; TURBT, transurethral resection of bladder tumor; ULN, upper limit of normal.

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Author Affiliations

1. Urology Associates, Nashville, TN; 2. Atlantic Urology Clinics, Myrtle Beach, SC; 3. University of California, San Diego, La Jolla, CA; 4. Tyra Biosciences, Inc., Carlsbad, CA; 5. Baylor College of Medicine, Houston, TX

Acknowledgments

The authors would like to thank all patients, caregivers, and all the SURF302 investigators and study personnel at study sites. Medical writing assistance was provided by Federica Rinaldi, PhD, of MEDI STRAVA (Yardley, PA), and was funded by Tyra Biosciences, Inc. This study was funded by Tyra Biosciences, Inc.

Sponsor Contact Information

TyraClinicalTrials@tyra.bio | Tyra Biosciences, Inc. | Carlsbad, CA, USA

Presented at the EAU26; March 13-16, 2026; London, England.

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