## Phase 2b Clinical Trial of IGV-001 for Patients With Newly Diagnosed Glioblastoma



## **ENROLLMENT COMPLETED**

## Phase 2b Trial to Further Assess the Safety and Efficacy of IGV-001 ClinicalTrials.gov identifier: NCT04485949

Phase 2 protocol [Lee IY, et al. Future Oncol 2024;20(10):579-591]



# PRIMARY OBJECTIVE

#### **SECONDARY OBJECTIVE** Survival overall

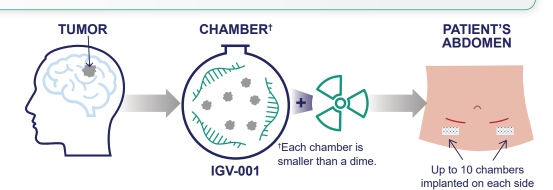
### **SAFETY OBJECTIVE**

Safety and tolerability

### **Treatment**

WEEK

IGV-001 is a personalized therapy that aims to induce antitumor immunity; it includes the patient's own glioblastoma tumor cells plus a molecule called IMV-001 placed in chamber



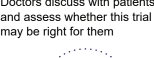
SCREENING Day -14 through Day -3

SCREENING/RANDOMIZATION Day -2 through Day 0

TREATMENT

Day 1 through Day 28

Doctors discuss with patients and assess whether this trial may be right for them



Patient with newly diagnosed glioblastoma (age 18-70 years)

MRI\*

Doctors will confirm brain tumor using frozen sample

Randomize 2:1 (N=93)

Superficial abdominal incision to the rectus muscle (after randomization) n=62 Patients are randomlv assigned to IGV-001 or placebo groups Placebo n=31

IGV-001

chambers chambers MRI\* within 3 days after

surgery

Day 3

remove

Tumor tissue will be used to prepare IGV-001; chambers will contain either IGV-001 or placebo



The usual treatment is RT\* and chemotherapy

RT\* + TMZ\*

Week 7-12

Surgery to remove tumor

**SOC\* TREATMENT** Week 7 through Week 41

FOLLOW-UP

Follow-up

Month 10-36

Month 10 through Month 36

Day 1

insert

Doctors keep track of patients' health after treatment

Days

14 and 28

examine

patients

IGV-001 was generally well tolerated

MRI\* 4 weeks (±1) after end of RT'

MRI\* 8 weeks (±2) after post RT\* MRI\*

TMZ\*

Week 17-41

MRI\* every 3 months (±2 weeks) until 36 months after randomization or disease progression

Phase 1b Trial Results [Andrews DW, et al. Clin Cancer Res. 2021;27(7):1912-1922]

had mild or non–life-threatening AEs\* related to the cut in the abdomen where the chambers were placed

had mild or non–life-threatening AEs that may have been caused by IGV-001

**EFFICACY** 

**SAFETY** 

Patients who received IGV-001 and SOC\* in past studies lived, on average, longer than those in earlier studies who received only SOC\*



**10.6 MONTHS LONGER** without worsening of disease



22 MONTHS LONGER overall

## FACTSHEET: Phase 2b Clinical Trial of IGV-001 for Patients With Newly Diagnosed Glioblastoma

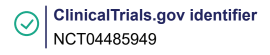


## ENROLLMENT COMPLETED

### **Protocol title**

A Randomized, Multicenter, Double-Blind, Placebo-Controlled, Phase 2b Study to Assess the Safety and Efficacy of IGV-001, an Autologous Cell Immunotherapy With Antisense Oligonucleotide (IMV-001) Targeting IGF-1R, in Newly Diagnosed Patients With Glioblastoma







## **Key Inclusion Criteria**



### Patients who take part in the trial\* must:

- Have newly diagnosed glioblastoma
- Be 18 to 70 years of age
- Have a KPS score ≥70 (unable to work but able to care for themselves overall)



## **Key Exclusion Criteria**





### Patients are not allowed to participate\* in the trial if they have:

- A tumor that is on both sides of the brain
- Had previous surgery or anticancer treatment for glioblastoma
- Glioblastoma that came back
- Another cancer<sup>†</sup> while having glioblastoma or within the last 3 years that is not cured
- A weakened immune system (example: HIV, HBV, HCV) or an autoimmune disorder (example: Crohn's disease)
- Heart disease or history of heart issues



SCREENING: Patients will have screening procedures completed between Day -14 to Day -2 (up to 16 days)

RANDOMIZATION: Patients are randomly assigned 2:1 to treatment with IGV-001 or placebo

TREATMENT: Patients receive study treatment (IGV-001 or placebo) during Days 1-28

**SOC TREATMENT:** Patients receive usual treatment (SOC) of RT and chemotherapy (TMZ) during Weeks 7-12, then chemotherapy alone during Weeks 17-41

FOLLOW-UP: Doctors keep track of patients' health during Months 10-36

<sup>\*</sup>Additional criteria apply. Please refer to protocol 14379-201 for full inclusion and exclusion criteria. †Patients can participate if they had some skin cancers, superficial bladder cancer (cancer that was only on the surface of the lining of the bladder), or carcinoma in situ (cancer that had not spread) of the cervix or breast