

# GENETIC TEST REPORT

## Patient Details:

Name: John Doe  
Date of birth: 18 March 1995  
Sex: Male  
NHS number: NH00198  
Sample type: Blood

## Test ordered by:

Name: Dr Requesta  
Organisation: Chester Hospital  
Telephone: 01223 555555  
Copies to: Dr A. Nother  
Dr X. Tra

## Test carried out by:

Laboratory: Gentest UK  
Telephone: 01223 666555  
Date received: 26 February 2018  
Date reported: 12 March 2018  
Authorised by: A Tester

## REASON FOR TEST

CF carrier status testing requested. Partner is a carrier of CF.

## YOUR RESULT: No common cystic fibrosis gene alterations detected

### ABOUT THE TEST

This test looked at a gene called CFTR. Everyone has two copies of this gene (one from their mother and one from their father). Alterations to this gene can cause the condition cystic fibrosis (CF).

If you have an alteration in **both** copies of your CFTR genes you will have CF.

If you have an alteration in **only one** copy of CFTR you will not have CF but will be a **'carrier'**.

Carriers are healthy but may pass on their altered gene to any children.

### WHAT THIS RESULT MEANS FOR YOU

Your sample has been tested for the common alterations to the CFTR gene that cause CF. No alterations were detected, so you are not a carrier of any of the alterations we tested for.

There are many alterations to the CFTR gene that are known to cause CF but some are very rare. This test looks for almost all of the alterations seen in the UK population.

Around 6 in 1000 people who receive this test result (0.6%) carry a CFTR gene alteration: there is still a very small possibility that you have one of the rare alterations. However, this is highly unlikely if there is no-one in your family who has CF.

If you have children with someone who is a carrier of CF, there is a less than 2 in 1000 (0.16%) chance in each pregnancy that the child will have CF.

### NEXT STEPS

- **You do not need to do anything as a result of this test, but if you have questions about it, talk to your doctor.**

### MORE INFORMATION AND SUPPORT

If you have questions about your test result, talk to the doctor who ordered your test. Your doctor can also phone or write to the East Anglian Clinical Genetics Service on 01223 216446 to make a referral or ask for further advice to help answer your queries.

To understand more about genetic testing, visit: [www.nhs.uk/conditions/genetics/services/](http://www.nhs.uk/conditions/genetics/services/)

To understand more about cystic fibrosis, visit: [www.cysticfibrosis.org.uk](http://www.cysticfibrosis.org.uk)

# FOR YOUR RECORDS

## Technical Information

### Patient Details:

Name: John Doe  
Date of birth: 18 March 1995      Sample type: Blood  
Sex: Male      Sample collected: 10 March 2018  
NHS number: NH00198      Sample received: 12 March 2018

The information on this page is for health professionals. It is not essential that patients read this section.



Gene	Zygoty	Inheritance	HGVS Description	Location	Classification
CFTR	No variant	Not applicable	NM_000492.3: c.1521_1523delCTT p.(Phe508del) GRCh37(hg19)	Not applicable	Not applicable

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## Full Interpretation

John Doe's Partner, Jane (d.o.b. 28/09/2004) is a carrier of cystic fibrosis (CF) and is heterozygous for the common c.1521\_1523delCTT p.(Phe508del) cystic fibrosis gene (CFTR) variant (information received with referral).

DNA from John Doe has been tested for the common c.1521\_1523delCTT p.(Phe508del) CFTR gene variant as well as other CFTR variants (see below) which together account for approximately 85% of UK CFTR gene variants (WHO, 2004). No CFTR gene variants were detected by our test.

This reduces John Doe's CF carrier risk to 1 in 161 (0.6%).

## Test Methodology

CF-EU2v1 Elucigene Assay (<http://www.elucigene.com/product/cfeu2v1/>) includes 50 of the most frequently observed European CF pathogenic variants - see our website (<http://cuh.org.uk/genetics-labs-cf>) for variant details. Variant nomenclature is in accordance with the Human Genome Variation Society (HGVS) guidelines according to GenBank accession number NM\_000492.3 (DNA) and NP\_000483.3 (protein) where +1 is the A of the ATG translation initiation codon. Very rare variants within the CFTR gene may interfere with the CFEU2v1 assay, causing false positive or false negative results. Please see the Elucigene website (<https://www.elucigene.com>) for further information regarding CFTR analysis.

## References:

World Health Organization (WHO), 2004. The molecular epidemiology of cystic fibrosis. Available: <http://www.who.int/genomics/publications/reports/en/>

McCormick et al. 2002 Eur J Hum Genet. 2002 Oct;10(10):583-90.