Revised HOME-CoV - Study on the implementation of the revised HOME-CoV score to guide the choice of hospitalisation or outpatient management of patients with confirmed or probable SARS-CoV-2 infection admitted to an emergency department.

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Address

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General			
Identification			
Detailed name	Study on the implementation of the revised HOME-CoV score to guide the choice of hospitalisation or outpatient management of patients with confirmed or probable SARS-CoV-2 infection admitted to an emergency department.		
Sign or acronym	Revised HOME-CoV		
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	2020-A03067-32		
General Aspects			
Medical area	Emergency medicine		
Study in connection with Covid- 19	Yes		
Health determinants	Healthcare system and access to health care services		
Keywords	COVID-19; Hospitalization; Outpatient; Rule validation; Expert consensus; Rule-based decision-making; Clinical support decision tool.		
Scientific investigator(s) (Contact)			
Name of the director	Douillet		
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Collaborations		
Funding		
Funding status	Public	
Details	Angers University Hospital	
Governance of the database		
Sponsor(s) or organisation(s) responsible	Angers University Hospital	
Organisation status		
Additional contact		
Main features		
Type of database		
Type of database	Study databases	
Study databases (details)	Cohort study	
Database recruitment is carried out by an intermediary	A selection of health institutions and services	
Database recruitment is is made on the basis of:	Another treatment or procedure	
Database recruitment is carried out as part of an interventional study	Yes	
Details	Performed at group level (clusters)	
Database objective		
Main objective	The primary objective is to demonstrate the reliability and safety of outpatient management among patients highly suspected or confirmed as infected with COVID-19, attending an emergency department and with a revised HOME-CoV score less than 2 (negative rule).	
	The secondary objectives are as follows: i. Evaluate the rate of patients having required hospitalisation within 7 days following inclusion according to the revised positive or negative HOME-	

CoV rule.

ii. Evaluate the rate of patients having required hospitalisation and initiation of oxygen therapy within 7 days following inclusion according to the revised positive or negative HOME-CoV rule.
iii. Evaluate the rate of patients having required intubation within 7 days following inclusion according to the revised positive or negative HOME-CoV rule.

iv. Evaluate the rate of all-cause deaths within 7 days following inclusion according to the revised positive or negative HOME-CoV rule.
v. Evaluate the performance of the revised HOME-CoV score in predicting a negative outcome in the patient subgroup with SARS-CoV-2 infection

confirmed by RT-PCR to rule out the risk of a negative outcome when assessed as low risk. vi. Compare the performance of the revised HOME-CoV score with that of other existing scores.

Inclusion criteria

Adult patient (> 18 years),

- Attending one of the emergency departments taking part in the study due to COVID-19 infection confirmed by SARS-CoV-2 positive RT-PCR, or considered highly probable by the physician managing the patient,
- Not requiring management in a continuous care or intensive care unit, and subject to a decision to limit active treatment,
- Having given their formal consent to take part in the study,
- Registered with or a beneficiary of a social security scheme.

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Age Adulthood (19 to 24 years)
Adulthood (25 to 44 years)

Adulthood (45 to 64 years) Elderly (65 to 79 years)

Great age (80 years and more)

Population covered Sick population

Pathology B33 - Other viral diseases, not elsewhere classified

Gender Male Woman

Woman

Geography area International

Detail of the geography area France Belgium

Data collection **Dates** Size of the database Size of the database (number of [1000-10 000[individuals individuals) Details of the number of 1,300 patients individuals Data Current data collection Database activity Type of data collected Clinical data Declarative data Paraclinical data Biological data Administrative data Clinical data (detail) Direct physical measures Medical registration Declarative data (detail) Face to face interview **Procedures** Followed pathology Promotion and access Promotion

Access