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1. ≥18 years

2.	ESKD on haemodialysis con peritoneal dialysis		CKD stage 4 or 5 (eGFR \leq 29 mL/min/ 1.73 m ²) not receiving renal replacement therapy	
	 Receiving maintenance haemodialysis or peritoneal dialysis for at least 90 days/ 3 months, <i>and</i> Irreversible kidney failure (opinion of the treating nephrologist). 		 eGFR <29 mL/min/1.73 m2 for >3 months, and Not currently receiving maintenance haemodialysis or peritoneal dialysis, and Not have a functioning kidney allograft in a kidney transplant recipient. 	

3.	Elevated CV r	isk, defined by at least one of :				
	History of CAD or	 One or more of: Myocardial infarction, or Multi-vessel PCI or CABG surgery, or Single-vessel PCI or CABG surgery and stenosis of greater than or equal to 50% in at least one other coronary artery, confirmed by invasive coronary angiography, or non-invasive imaging or stress studies (e.g. exercise or pharmacologic) suggestive of significant ischemia, or Medically managed multi-vessel coronary disease with symptoms or with history of stable or unstable angina. 	or	Diabetes mellitus	or	≥65 years
	PAD	 One or more of : Previous aorto-femoral bypass surgery, limb bypass surgery, or percutaneous transluminal angioplasty revascularization of the iliac, or infra-inguinal arteries, or Previous limb or foot amputation for arterial vascular disease, or History of intermittent claudication and one or more of the following: An ankle/arm BP ratio <0.90, or Significant peripheral artery stenosis (≥50%) documented by angiography, or by duplex ultrasound, or Previous carotid revascularization or asymptomatic carotid artery stenosis ≥50% as diagnosed by duplex ultrasound or angiography 				
	Non- haemorrhagic non- lacunar stroke	 One or more of: Non-haemorrhagic non-lacunar stroke more than one month prior to study enrolment, or Diffusion-weighted imaging (DWI) positive TIA on magnetic resonance imaging more than one month prior to study enrolment. 				

Exclusion Criteria





 Heart valve, Indication/contraindication to 	10. Severe heart failure with ejection fraction <30% or NYHA class III or IV symptoms,				
anticoagulant therapy,	11. Hypersensitivity or contraindication				
3. High bleeding risk/coagulopathy,	to rivaroxaban,				
 Lesion/condition of significant risk of major bleeding*, 	12. Uncontrolled hypertension (≥180/110 mm Hg) at screening,				
5. Major bleeding episode within 30 days prior to study enrolment, or active and clinically	13. Haemoglobin <90g/L, or platelet count <100 x 109/L,				
significant bleeding, 6. P2Y12 inhibitors/adenosine diphosphate (ADP)	14. Significant liver disease or ALT >3 times upper normal limit.				
receptor inhibitors or phosphodiesterase inhibitors, physician/patient does not wish to stop medications,	15. Kidney transplant recipients with a functioning allograft, or scheduled for living-donor kidney transplant surgery,				
7. Strong inhibitors of combined CYP3A4 and P-glycoprotein; or strong inducers of CYP3A4,	16. Pregnancy/ intention to become pregnant/ breast-feeding,				
8. Stroke within 1 month,	17. Inability to understand or comply with the				
9. History of a haemorrhagic or lacunar stroke,	requirements of the study.				
* Examples for risk of major bleeding					
gastrointestinal ulceration,	• arteriovenous malformations (excluding AV fistula				
• malignant neoplasms at high risk of bleeding,	or AV graft for dialysis vascular access), • vascular aneurysms or major intraspinal or intracerebral vascular abnormalities, • branchiastasis or pulmonary blanding				
• brain or spinal injury,					
 brain or spinal or ophthalmic surgery, 					
intracranial haemorrhage,	 bronchiectasis or pulmonary bleeding, congenital or acquired bleeding disorder. 				
• oesophageal varices,					

List of prohibited medications for TRACK eligibility

- Oral or parenteral anticoagulant treatment except for regional anticoagulation for haemodialysis
- P2Y12 inhibitors/ADP receptor inhibitors: clopidogrel, prasugrel, ticagrelor, cangrelor
- Phosphodiesterase inhibitor: dipyridamole
- Strong inhibitors of combined CYP3A4 and P-glycoprotein: » synthetic azole antimycotics, eg. ketoconazole, fluconazole, itraconazole, voriconazole, or posaconazole, if used systemically
 - »HIV-protease inhibitors, eg. ritonavir
 - » clarithromycin, erythromycin.
- Strong inducers of CYP3A4: » rifampicin, rifabutin, phenobarbital, phenytoin, carbamazepine, St John's wort.





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★ Contact National Lead or Chief Investigators if SUSAR is suspected

SAEs that are study outcomes

Primary efficacy outcomes

- » cardiovascular death
- » non-fatal myocardial infarction
- » stroke
- » peripheral artery disease event

Secondary and tertiary efficacy outcomes

- » All-cause death (including non-cardiovascular death and death due to undetermined cause)
- » venous thromboembolism
- » thrombosis of dialysis vascular access among participants with an AV fistula/graft

Safety outcomes

- » Major bleeding events, including fatal bleeding, symptomatic bleeding in a critical area or organ, bleeding leading to hospitalization,
- » Gastrointestinal bleeding

SAEs that are consistent with the natural history of advanced CKD/ESKD and associated conditions

- » Planned hospitalisations (for example, surgery, respite care, etc)
- » SAEs (including unplanned hospitalisations) that are expected to occur in high frequency in the study population.

Event	Report on	Timelines		
SAEs that are study outcomes	Outcome eCRF	Within 7 days of discharge from hospital		
SAEs consistent with natural history of advanced CKD/ESKD and associated conditions; planned admissions	SAE eCRF	Within 7 days of discharge from hospital		
Suspected unexpected serious adverse reactions (SUSARs)	SAE eCRF	Within 24 hours		
Any event of particular concern to the investigator*	SAE eCRF	Within 24 hours		
SAEs that are none of the above	SAE eCRF	Within 7 days of discharge from hospital		

^{*} Includes pregnancies





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