#### **PIET DEVELOPMENT OF AN IN-HOUSE METHOD FOR THE** Abstract ID #268 SYNTHESIS OF SODIUM [<sup>18</sup>F]FLUORIDE PREPARATION

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## Background and goal of the present work

Sodium [18F]fluoride is PET radiopharmaceutical containing fluorine-18 in the form of sodium fluoride, for visualization of the skeletal system and microcalcification.

Aim: Development of an in-house method for [18F]NaF production using only a dispensing module for both, radiopharmaceutical synthesis and dose dispensing.



### **Materials and methods**

For the production of [<sup>18</sup>F]NaF, the modification of the kit was done. On the dispensing kit, the Y connector and anion-exchange cartridge and sterile vial with 0.9% sodium chloride solution were installed. The preconditioned QMA cartridge was connected to the distal part of the Y connector and attached to the first valve of the kit. One proximal ending of the Y connector was connected to the [<sup>18</sup>F]F<sup>-</sup> cyclotron transfer line and the other proximal ending to the vial with 5 mL 0.9% NaCl solution for elution.





Semi-	The kit includes four three-way
automatic	valves, lines for different purposes,
dispenser	and a 0.22 µm sterilization filter

2102	
Recovery water	

### **Modified kit for production of [18F]NaF**



After development of our in-house method for synthesis, the process optimization was carried out, 10 production batches were performed, with yield higher than 98%, decay-corrected. All obtained QC results confirmed that the quality of [<sup>18</sup>F]NaF meets the acceptance criteria defined in the specification.

Test	Method	Acceptance criteria	Results	■1 900 2 µS □	St NaF 1mg/ml in HPLC water 04.10.2021 0h NaF0410211 0h	ECD_1 A2D_Analog	<1 250 _µS _	it NaF 1mg/ml in HPLC water 04.10.2021 0h NaF0410211 0h	ECD_1 ECD_1
Pre-release tests			800-	1 - 18F-Fluoride - 6.027		220	1 - Fluoride - 6.473 ∖\		
Appearance	Visual inspection	Clear, colorless solution	Clear colorless solution	700-			200		
Half-life Identification	Radioactivity measurements	1.75-1.92 hours	1,832 ± 0,014	600-			160 160		
Difference in retention time	HPLC	≤ 40 s	26,899 ± 4,763	500-			140- 120-		
Approximate pH value	pH strips	5.5 - 8.0	6.5-7.0				100		
Chemical purity: fluoride (F <sup>-</sup> )	HPLC	≤ 0.452 mg/mL	≤ 0.452				80		
Radiochemical purity: [ <sup>18</sup> F]fluoride	HPLC/gamma detector	min 98.5 % of the total radioactivity	100%		1 - Fluoride - 6.473		60- - - 40-		
Post-release tests									
Bacterial endotoxins	Chromogenic LAL method	≤ 17.5 IU/mL	< 5.00	 100 _2					
Sterility	Test for sterility (Ph. Eur)	Sterile	Sterile				-20-		
Radionuclidic purity	Gamma-ray	min 99.9 % of the	1,34 x 10 <sup>-05</sup> ±		4.0 6.0 8.0 10.0	min 12.0 15.0	-50	6.0 8.0 10.0 12	min .0 15.0
Fluorine-18	spectrometry	total radioactivity	7,63 x 10 <sup>-06</sup>		Padiachamical nurif			homical nurity	



# Conclusion

The in-house method for [<sup>18</sup>F]NaF synthesis was successfully designed and developed. The produced [<sup>18</sup>F]NaF radiopharmaceutical meets the acceptance criteria defined in the specification, based on Ph. Eur. monograph.

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