

Quels vaccins pour demain?

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Team GIMAP, CIRI, Inserm, U1111, CNRS, UMR530

Chaire Prévention, Vaccination, Contrôle de l'Infection PRESAGE

Déclaration de liens d'intérêt avec les industries de santé en rapport avec le thème de la présentation (loi du 04/03/2002) :

Intervenant : **Elisabeth Botelho-Nevers**

Titre : **Quels vaccins pour demain?**

L'orateur ne souhaite pas répondre

Consultant ou membre d'un conseil scientifique: **PAS de rémunération à titre personnel** (Pfizer, Janssen, Sanofi Pasteur)

OUI NON

Conférencier ou auteur/rédacteur rémunéré d'articles ou documents
PAS de rémunération à titre personnel

OUI NON

Prise en charge de frais de voyage, d'hébergement ou d'inscription à des congrès ou autres manifestations (**y compris pour cette présentation**)

OUI NON

Investigateur principal d'une recherche ou d'une étude clinique (**CIC**) (Sanofi Pasteur, GSK, Pfizer, MSD, Janssen, Moderna)

OUI NON

Membre du copil du I-REIVAC

22^{es} JNI, Montpellier du 30/08 au 1^{er}/09/2021

Petit préambule

- Intérêt des vaccins, globalement n'est plus à démontrer
- Populations changent, risques changent, émergences... bref toujours de nouveaux besoins!
- Prévenir vaudra toujours mieux que guérir

- Vaccination de masse
- Vaccination personnalisée
- La crise COVID a catalysé le changement et le développement vaccinal!

Pour demain: des vaccins meilleurs

Vaccin contre la tuberculose

- La tuberculose reste dans le monde une des principales causes de mortalité infectieuse: 1,4 millions de décès dont 210 000 PVVIH en 2019
- ≈ 1,7 milliards de personnes infectées, 5-15% évolueront vers une tuberculose maladie: ID, âge avancé...
- BCG: CI chez ID, durée de l'immunité variable (10-20 ans, max 60), meilleure efficacité si IDR négative, efficacité modérée (0-77%!)
- Besoin de nouveaux vaccins

Prévention de la maladie tuberculeuse

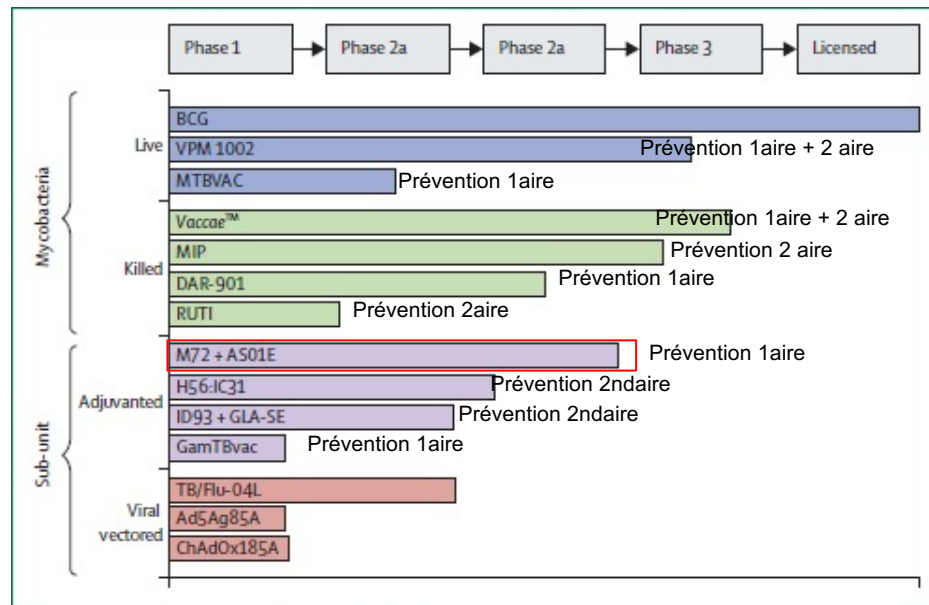


Figure: Tuberculosis vaccine candidates in clinical development

The indicated clinical development stages of vaccine candidates are based on an extrapolation from data in ClinicalTrials.gov.

Rapport sur la tuberculose dans le monde 2020 : résumé d'orientation OMS, [http://apps.who.int/iris/Schrager_LK, et al., Lancet Infect Dis. 2020 Mar;20\(3\):e28-e37](http://apps.who.int/iris/Schrager_LK_et_al.,_Lancet_Infect_Dis._2020_Mar;20(3):e28-e37).
Brazier b et al., Seminars in Immunopathology (2020) 42:315–331
<https://www.clinicaltrials.gov>

Vaccin contre la tuberculose

The NEW ENGLAND JOURNAL of MEDICINE

N Engl J Med 2018; 379: 138–49.

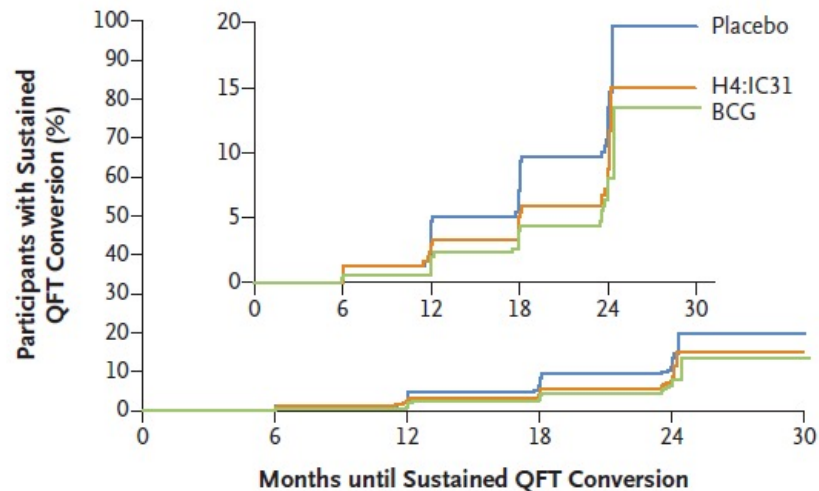
ORIGINAL ARTICLE

Prevention of *M. tuberculosis* Infection with H4:IC31 Vaccine or BCG Revaccination

E. Nemes, H. Geldenhuys, V. Rozot, K.T. Rutkowski, F. Ratangee, N. Bilek, S. Mabwe, L. Makhethe, M. Erasmus, A. Toefy, H. Mulenga, W.A. Hanekom, S.G. Self, L.-G. Bekker, R. Ryall,* S. Gurunathan, C.A. DiazGranados, P. Andersen, I. Kromann, T. Evans, R.D. Ellis, B. Landry, D.A. Hokey, R. Hopkins, A.M. Ginsberg, T.J. Scriba, and M. Hatherill, for the C-040-404 Study Team†

the BCG vaccine reduced the rate of sustained QFT conversion, with an efficacy of 45.4% ($P = 0.03$); the efficacy of the H4:IC31 vaccine was 30.5% ($P = 0.16$).

C Sustained QFT Conversion



No. at Risk

	0	6	12	18	24	30
Placebo	310	302	287	263	122	
H4:IC31	308	303	288	268	124	
BCG	312	308	297	281	136	

Vaccin contre la tuberculose

The NEW ENGLAND JOURNAL of MEDICINE

GSK

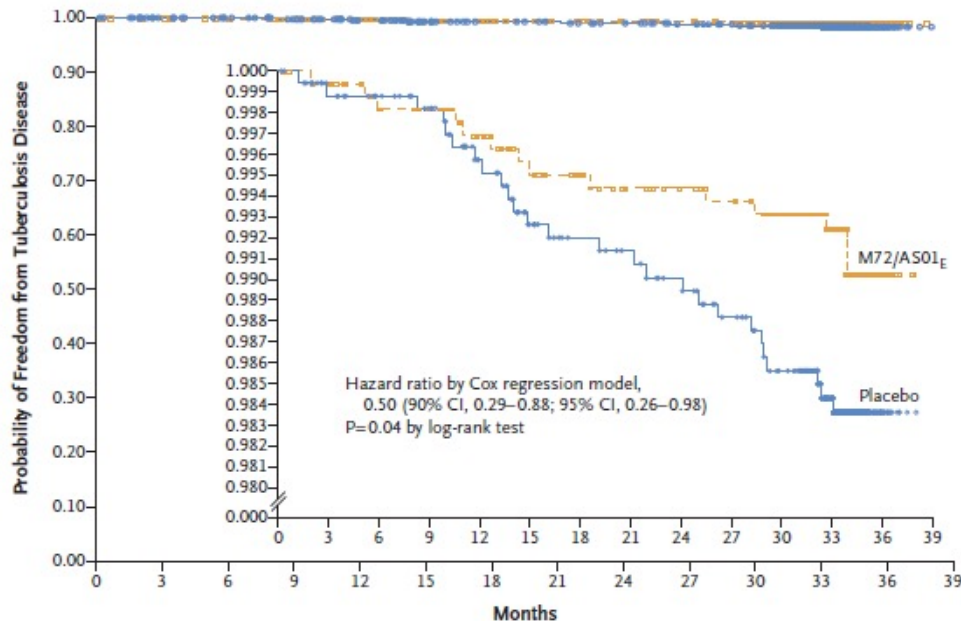
N Engl J Med 2019; 381: 2429-2439

ORIGINAL ARTICLE

Final Analysis of a Trial of M72/AS01_E Vaccine to Prevent Tuberculosis

D.R. Tait, M. Hatherill, O. Van Der Meeren, A.M. Ginsberg, E. Van Brakel, B. Salaun, T.J. Scriba, E.J. Akite, H.M. Ayles, A. Bollaerts, M.-A. Demoiitié, A. Diacon, T.G. Evans, P. Gillard, E. Hellström, J.C. Innes, M. Lempicki, M. Malahleha, N. Martinson, D. Mesia Vela, M. Muyoyeta, V. Nduba, T.G. Pascal, M. Tameris, F. Thienemann, R.J. Wilkinson, and F. Roman

- M72: protéine de fusion
- 08/2014 à 11/2015, inclusion adultes 18 -50 ans IGRA+, VIH -, Sans évidence de tuberculose maladie
- Kenya, Afrique du Sud et Zambie.
- Endpoint= Tuberculose maladie: PCR+, culture crachats
- 3575 participants ont été randomisés 1:1, 3573 ont reçu au moins une dose de M72/AS01E ou de placebo, et 3330 ont reçu les 2 doses.
- 2 doses à 1 mois d'intervalle, suivi 3 ans



No. at Risk

	0	3	6	9	12	15	18	21	24	27	30	33	36	39
M72/AS01 _E	1626	1621	1614	1609	1592	1580	1573	1566	1561	1557	1542	1468	12	
Placebo	1663	1650	1642	1632	1610	1586	1576	1571	1564	1553	1539	1460	19	

Figure 1. Kaplan-Meier Estimate of Definite Pulmonary Tuberculosis According to the First Case Definition.

Profil de sécurité tout à fait correct
Essai programmé chez les PVVIH (NCT04556981)

Vaccin anti-pneumococcique

Invasive pneumococcal disease incidence in children and adults in France during the pneumococcal conjugate vaccine era: an interrupted time-series analysis of data from a 17-year national prospective surveillance study



Lancet Infect Dis 2020;
21: 137–47

Naim Ouldali, Emmanuelle Varon, Corinne Levy, François Angoulvant, Scarlett Georges, Marie-Cécile Ploy, Marie Kempf, Julie Creminier, Robert Cohen, Daniel Levy Bruhl*, Kostas Danis*

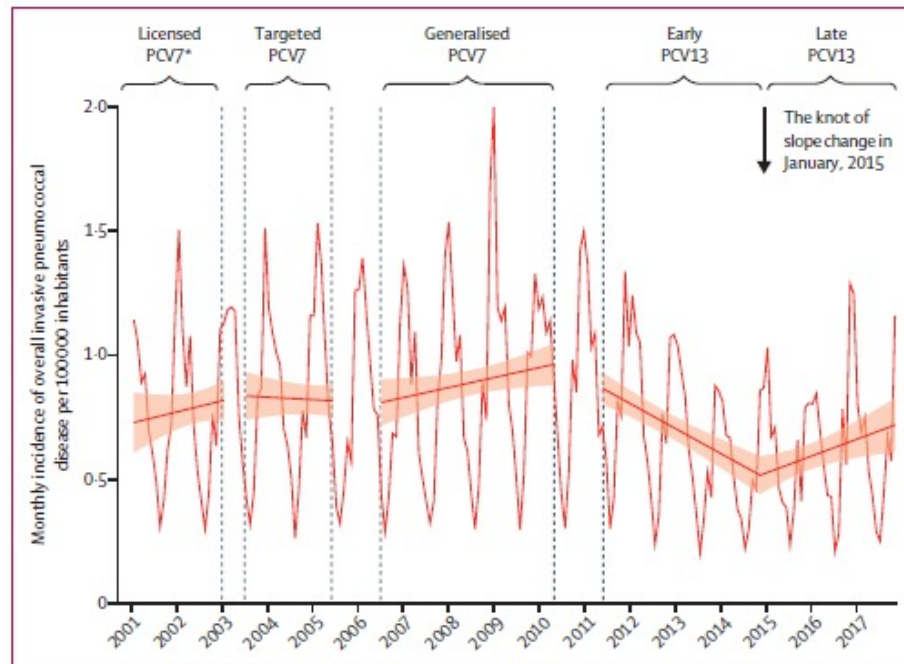


Figure 1: Time-series analysis of invasive pneumococcal disease incidence over 17 years

This figure represents data from 75 903 invasive pneumococcal disease cases. The bold slope lines were estimated by the segmented regression model; the red shading shows the 95% CI. The dotted vertical lines demarcate transition periods during which a new vaccine was implemented or changes to vaccination policy occurred. PCV=pneumococcal conjugate vaccine. Licensed PCV7=period from January, 2001, to December, 2002. Targeted PCV7=period from June, 2003, to May, 2005. Generalised PCV7=period from June, 2006, to May, 2010. Early PCV13=period from June, 2011, to December, 2014. Late PCV13=period from January, 2015, to December, 2017. *Licensed but not reimbursed PCV7 (vaccine coverage <10%).

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- Echappement/glisement sérotypes non contenus dans le PCV13
- Certains contenus dans le PPV23 mais pas tous, notamment 24F
- Schéma vaccinal de l'enfant ne contient pas le PPV23

* Sérotype contenu dans le PPV 23



	Licensed PCV7 period (January, 2001–December, 2002)	Targeted PCV7 period (June, 2003–May, 2005)	Generalised PCV7 period (June, 2006–May, 2010)	Early PCV13 period (June, 2011–December, 2014)	Late PCV13 period (January, 2015–December, 2017)
Children <2 years					
PCV7 serotypes (n=868)	67.9%	49.6%	9.8%	4.5%	5.8%
Serotypes specific to PCV13 plus serotype 6C (n=1187)	19.4%	31.6%	59.7%	15.1%	9.3%
Serotype 19A	9.3%	14.3%	29.1%	6.8%	3.7%
Serotype 3	3.2%	3.6%	4.2%	3.0%	4.7%
Main non-PCV13 serotypes (n=922)*	5.3%	10.4%	17.3%	55.2%	58.3%
Serotype 24F	1.5%	2.5%	5.7%	20.4%	24.4%
Serotype 15B/C *	1.8%	2.8%	3.1%	8.3%	8.1%
Serotype 10A *	0.4%	1.1%	2.4%	6.6%	6.4%
Serotype 12F *	0.2%	0.0%	1.7%	9.0%	4.8%
Serotype 22F *	0.2%	0.5%	1.9%	4.3%	4.8%
Serotype 8 *	0.6%	0.4%	0.7%	1.0%	4.0%
Serotype 15A	0.4%	2.0%	1.5%	4.7%	3.1%
Serotype 9N	0.2%	1.1%	0.4%	1.0%	2.9%
Adults ≥65 years					
PCV7 serotypes (n=2033)	50.8%	46.8%	23.2%	8.6%	6.6%
Serotypes specific to PCV13 plus serotype 6C (n=2601)	23.6%	27.8%	42.3%	36.9%	26.7%
Serotype 19A	8.7%	7.5%	14.6%	12.4%	7.4%
Serotype 3	8.2%	10.5%	10.1%	10.8%	14.3%
Main non-PCV13 serotypes (n=1977)	11.0%	11.8%	18.8%	30.3%	37.5%
Serotype 22F *	1.9%	3.5%	4.7%	6.8%	8.1%
Serotype 8 *	2.6%	1.6%	2.1%	2.3%	6.9%
Serotype 9N	1.7%	1.8%	2.4%	3.0%	5.3%
Serotype 12F *	0.2%	0.3%	2.1%	6.5%	4.8%
Serotype 15A	1.1%	0.8%	2.9%	5.1%	4.3%
Serotype 10A *	1.6%	0.9%	1.0%	2.5%	3.0%
Serotype 24F	1.4%	1.7%	2.7%	2.4%	2.7%
Serotype 15B/C *	0.5%	1.3%	0.9%	1.7%	2.4%

Vaccin anti-pneumococcique



Review

Development of Next Generation *Streptococcus pneumoniae* Vaccines Conferring Broad Protection

Malihe Masomian ¹, Zuleeza Ahmad ¹, Lai Ti Gew ²  and Chit Laa Poh ^{1,*} 

- ¹ Centre for Virus and Vaccine Research, School of Science and Technology, Sunway University, Kuala Lumpur, Selangor 47500, Malaysia; malihem@sunway.edu.my (M.M.); zuleezaa@sunway.edu.my (Z.A.)
² Department of Biological Sciences, School of Science and Technology, Sunway University, Kuala Lumpur, Selangor 47500, Malaysia; janeg@sunway.edu.my
* Correspondence: pohcl@sunway.edu.my; Tel.: +60-3-7491-8622 (ext. 7338); Fax: +60-3-5635-8633

Vaccines 2020, 8, 132; doi:10.3390/vaccines8010132

- PCV 15 (V114) Merck phase 3 (serotypes PCV13 + 22F et 33F)
- PCV 20 Pfizer (phase 3)
- *S. pneumoniae* killed whole-cell vaccine (WCV) (phase 2)
- PnuBioVax (*S. pneumoniae* serotype 4 TIGR4) (phase 1)
- PPrV (recombinant proteins, PcpA, PhtD, and PlyD1) Sanofi Pasteur (phase 2 en association avec PHiD-CV)

Vaccin anti-pneumococcique

Clinical Infectious Diseases

Clin Inf Dis 2020



MAJOR ARTICLE

Safety, Tolerability, and Immunogenicity of a 20-Valent Pneumococcal Conjugate Vaccine (PCV20) in Adults 60 to 64 Years of Age

Donald Hurley,¹ Carl Griffin,² Mariano Young Jr,² Daniel A. Scott,³ Michael W. Pride,⁴ Ingrid L. Scully,⁴ John Ginis,³ Joseph Severs,⁴ Kathrin U. Jansen,⁴ William C. Gruber,² and Wendy Watson³

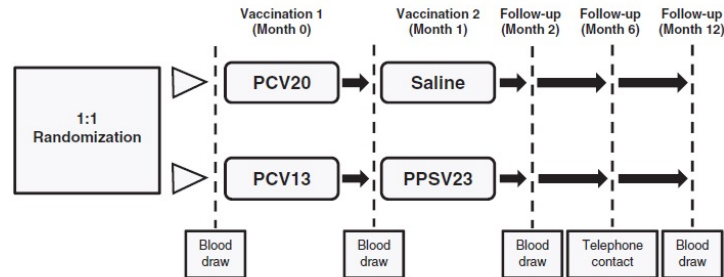
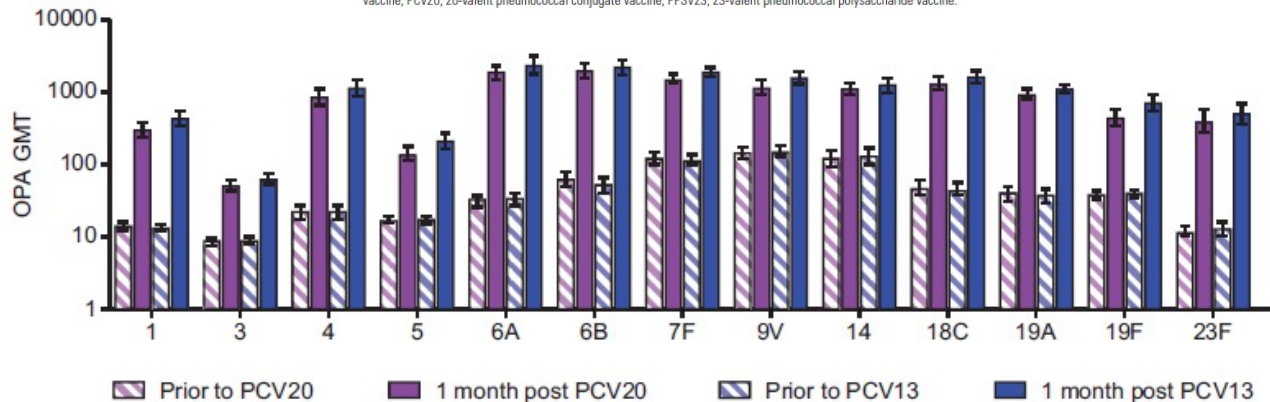


Figure 1. Study design. Immune responses measured at 12 months following vaccination 1 are not reported here. Abbreviations: PCV13, 13-valent pneumococcal conjugate vaccine; PCV20, 20-valent pneumococcal conjugate vaccine; PPSV23, 23-valent pneumococcal polysaccharide vaccine.

PCV 20=PCV 13+ nouveaux serotypes (8, 10A, 11A, 12F, 15B, 22F, and 33F)

PPV23 sert de contrôle pour l'immunogénicité des 7 serotypes additionnels du PCV20 (8, 10A, 11A, 12F, 15B, 22F, 33F).



GMFRs in Functional Antibody From Baseline 1 Month After Vaccination

Serotype	1	3	4	5	6A	6B	7F	9V	14	18C	19A	19F	23F
PCV20	21.1	6.1	37.1	8.2	57.4	29.0	12.3	7.7	8.3	26.4	22.6	11.5	32.9
PCV13	33.5	7.1	51.0	11.6	68.6	38.8	15.8	10.1	9.6	35.2	30.9	18.4	39.8



22^{es} JNI, Montpellier du 30/08 au 1^{er}/09/2021

Vaccin anti-pneumococcique

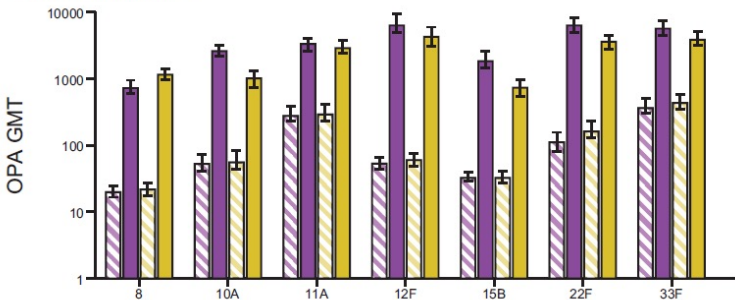
Clinical Infectious Diseases

MAJOR ARTICLE



Safety, Tolerability, and Immunogenicity of a 20-Valent Pneumococcal Conjugate Vaccine (PCV20) in Adults 60 to 64 Years of Age

Donald Hurley,¹ Carl Griffin,² Mariano Young Jr,² Daniel A. Scott,³ Michael W. Pride,⁴ Ingrid L. Scully,⁴ John Ginis,³ Joseph Severs,⁴ Kathrin U. Jansen,⁴ William C. Gruber,⁴ and Wendy Watson¹



Prior to PCV20 1 month post PCV20
 Prior to PPSV23 1 month post PPSV23

GMFRs in Functional Antibody From Baseline 1 Month After Vaccination

Serotype	8	10A	11A	12F	15B	22F	33F
PCV20	36.7	47.5	11.0	112.2	56.7	54.4	14.0
PPSV23	56.4	17.0	9.7	76.1	20.8	20.0	9.0

Table 2. Summary of Adverse Events (Safety Population)

Time Point Type of AE	PCV20/Saline (n = 221 ^a /213 ^b)		PCV13/PPSV23 (n = 222 ^a /214 ^b)	
	n (%)	(95% CI)	n (%)	(95% CI)
Following PCV20 or PCV13 administration through 1 month of follow-up				
Any AE	27 (12.2)	(8.2, 17.3)	29 (13.1)	(8.9, 18.2)
Severe AE	3 (1.4)	(.3, 3.9)	3 (1.4)	(.3, 3.9)
SAE	0	(.0, 1.7)	1 (0.5)	(.0, 2.5)
NDCMC	2 (0.9)	(.1, 3.2)	1 (0.5)	(.0, 2.5)
Following saline or PPSV23 administration through 1 month of follow-up				
Any AE	15 (7.0)	(4.0, 11.3)	40 (18.7)	(13.7, 24.6)
Severe AE	1 (0.5)	(.0, 2.6)	6 (2.8)	(1.0, 6.0)
SAE	0	(.0, 1.7)	4 (1.9)	(.5, 4.7)
NDCMC	2 (0.9)	(.1, 3.4)	4 (1.9)	(.5, 4.7)
From 1 month following saline or PPSV23 administration through 12 months of follow-up				
SAE	9 (4.2)	(2.0, 7.9)	7 (3.3)	(1.3, 6.6)
NDCMC	9 (4.2)	(2.0, 7.9)	3 (1.4)	(.3, 4.0)
Throughout the study				
SAE	9 (4.1)	(1.9, 7.6)	11 (5.0)	(2.5, 8.7)
NDCMC	13 (5.9)	(3.2, 9.8)	8 (3.6)	(1.6, 7.0)

Poursuite en phase 3
Ne couvre toujours pas le 24F



22^{es} JNI, Montpellier du 30/08 au 1^{er}/09/2021

Vaccin antigrippal

- Efficacité vaccinale des vaccins inactivés faible, variable selon les années, populations (10-60%)
- Amélioration via fortes doses, méthode d'administration, des adjuvants
- Durée protection courte
- Toujours souche dépendante+++

TABLE 1 Advantages and disadvantages of different influenza vaccines

Vaccine Type	Advantage	Disadvantage
Inactive vaccine	a) All age groups (except children under 6 months) with no contraindications can receive inactivated influenza vaccine b) Safe in pregnant women c) Use in immunocompromised patients	a) Soreness at the vaccination site, fever, headache, myalgia, or any physical unease happen mostly in children b) Allergy c) In rare case autoimmune disorders
Live attenuate	a) Safe in cystic fibrosis patients b) No systemic allergic reactions such as urticaria, angioedema, rhinitis, and eczema	a) Mild to moderate symptoms including runny nose, sneezing, nasal discomfort, fever and headache b) Is not recommended to be routinely used in pregnant women
Recombinant	a) High safety profile without involving infectious viruses b) Rapid, stable c) Induces humoral and cellular immune responses	a) Low immunogenicity b) Require appropriate adjuvants
DNA vaccine	a) Induce all three arms of adaptive immunity, CTLs, antibodies, and helper T cells b) Possible mucosal delivery and thus may stimulate innate immunity	a) Lower immunogenicity, low level of T-cell, and B-cell memory due to b) Integration of DNA vaccine genetic material into cellular or host DNA, c) Development of autoimmune disorders against host DNA
Universal vaccine	M2e: a) Induces M2e-specific humoral and cellular immune responses; b) Elicits broad cross-protection against divergent virus strains Epitope-base: a) They are considered to be safe, easy to produce, and stable. b) Can induce B-cell and T-cell in the same formulation	a) Single M2e molecule induces lower immune responses a) The main disadvantage of the epitope-based vaccine is that algorithms may fail to predict all the appropriate epitopes
CTL inducing vaccine	a) Target conserved influenza virus proteins and improve recovery and inhibit disease progression	a) Need to have an epitope that can be recognized by all major histocompatibility complex (MHC)
RNA vaccine	a) Safety b) Efficacy c) Higher potency (especially with self-amplifying RNA vaccines)	a) Possibility of adverse consequences like thrombus and/or edema b) Limited availability in cases of pandemic and endemic diseases.

Vaccin antigrippal

- Task force: développement vaccin antigrippal universel
- Cibles: HA stalk protein (stable), NA, matrix protein 2 (M2) et la nucleoprotéine (NP).
- M2 et NP sont conservées au sein des souches humaines et aviaires

Universal influenza vaccine candidates in clinical development						
Targeted response	Vaccine target	Vaccine platform	Phase	Candidate name	Development partners	
B cell (antibody) responses to conserved regions of the virus	NA, HA gene suppression HA	LAIV	1	CodaVax	Codagenix, Inc. (US)	
		mRNA	1	Modified mRNA lipid nanoparticles H1ssF_3928	Moderna, Inc. (US)	
	HA stalk HA stalk, HA head	Ferritin-based nanoparticles	1		NIAID Vaccine Research Center (US)	
		Recombinant HA	3	Nano-Flu	Novavax (US)	
		Recombinant subunit VLP	1	ACAM-FLU-A	Sanofi Pasteur (US)	
		Viral vector	2	VXA-A1.1	Vaxart, Inc. (US)	
Cross-protective T cell responses against the virus' internal proteins	M2e	Recombinant fusion protein	1	Vax102	VaxInnate Corp (US)	
	NP, M1, PB1, PB2	Synthetic peptide	1	FP-01.1	Altimmune (US) (Immune Targeting Systems Ltd)	
		LAIV	2	M2SR	Flugen, Inc. (US)	
		Synthetic peptide	2	FLU-v	Imutex Ltd (SEEK/hVIVO) (UK)	
	B and T cell responses	NP, M1	Viral vector	1	MVA/ ChAdOx1-NP + M1	Jenner Institute/University of Oxford (UK)
		NP	Nanoparticles	2	OVX836	Oxivax SAS (France)
			Viral vector	2	MVA-NP + M1	Vaccitech (UK)
		HA (H1)	Viral vector	2	NasoVAX	Altimmune, Inc. (US)
Recombinant peptide			3	Multimeric-001	BlondVax Pharmaceuticals (Israel); NIAID; Seventh Framework Program (EU)	
NP, M1, HA2		Fusion protein	1	N8205	Dynavax (US)	
		DNA	1	VGX-3400	GeneOne Life Sciences, Inc. (South Korea)	
NA, cHA, HA head, HA stalk	NA, cHA, HA head, HA stalk	Functional cHA	1	Chimeric HA (cHA)-based vaccines	Mount Sinai School of Medicine (US); GSK (US); PATH (US)	
	NP, NA, HA	DNA	1	INO-4301	Inovio Pharmaceuticals (US)	
		VLP	3	Quadrivalent VLP (QVLP)	Medicago, Inc. (Canada)	
	M2e, HA2 stalk epitopes NS1-deficient	Recombinant protein	1	Uniflu	VA Pharma LLC (Russia); Russian Federation Ministry of Health	
LAIV		1	deltaFLU	Vivaldi Biosciences (US); Icahn School of Medicine at Mount Sinai (US); AVIR Green Hills Biotechnology AG (Austria)		

Source: CIDRAP, Universal Influenza Vaccine Technology Landscape; URL: <http://www.cidrap.umn.edu/universal-influenza-vaccine-technology-landscape>

Abbreviations: Chimeric HA (cHA); Hemagglutinin (HA); Influenza B virus (IBV); Live attenuated influenza virus vaccine (LAIV); Matrix Protein (M1); Membrane protein (M2); Membrane protein ion channel ectodomain (M2e); Neuraminidase (NA); Nonstructural protein (NS1); Nucleoprotein (NP); Viral RNA polymerases (PB1, PB2); Virus-like particle (VLP).

Vaccin anti-coronavirus

Science

A universal coronavirus vaccine

Wayne C. Koff and Seth F. Berkley

Science 371 (6531), 759.
DOI: 10.1126/science.abh0447

Vaccine 39 (2021) 4239–4241



Contents lists available at ScienceDirect

Vaccine

journal homepage: www.elsevier.com/locate/vaccine



Commentary

The need for broadly protective COVID-19 vaccines: Beyond S-only approaches

Gregory A. Poland*, Inna G. Ovsyannikova, Richard B. Kennedy

Mayo Clinic Vaccine Research Group, Mayo Clinic, Rochester, MN, USA



CellPress

Cell

Article

A Universal Design of Betacoronavirus Vaccines against COVID-19, MERS, and SARS

Lianpan Dai,^{1,2,3,12,*} Tianyi Zheng,^{1,2,12} Kun Xu,^{3,12} Yuxuan Han,^{2,12} Lili Xu,^{4,12} Enqi Huang,⁵ Yaling An,¹ Yingjie Cheng,⁵ Shihua Li,⁶ Mei Liu,⁷ Mi Yang,⁷ Yan Li,⁶ Huijun Cheng,¹ Yuan Yuan,⁶ Wei Zhang,⁶ Changwen Ke,⁸ Gary Wong,^{9,10} Jianxun Qi,^{2,6} Chuan Qin,^{4,*} Jinghua Yan,^{6,7,*} and George F. Gao^{1,2,6,11,13,*}



Pour demain: des vaccins nouveaux

Vaccin contre VIH

- En 2019, 38 millions de personnes vivent avec le VIH au niveau mondial, 1,7 millions de nouvelles infections
- Echec de nombreux essais vaccins depuis > 30 ans
- Echec récent de la phase 3 de l'essai HVTN702 (6ème essai clinique complet)

Recent efficacy trials and their related phase 1/2a trials.

Immunogen	NCT Trial number	Other names	Phase	Adjuvant	Completion Date*
Vector: ALVAC-HIV (vCP2438), Protein: bivalent subtype C gp120	NCT03284710	HVTN107	1/2a	Alum vs. MF59	Dec 2019
Vector: ALVAC-HIV (vCP2438), Protein: bivalent subtype C gp120	NCT03122223	HVTN120	1/2a	MF59 vs. AS01 _g	Jul 2020
Vector: ALVAC-HIV (vCP2438), Protein: bivalent subtype C gp120	NCT02968849	HVTN702/Uhambo	2b/3	MF59	Aug 2021; prematurely terminated
Vector: Ad26.Mos4.HIV, Protein: subtype C gp140 and/or mosaic gp140	NCT02935686	ASCENT/HVTN118/HPX2003	1/2a	Alum	Jan 2022
Vector: Ad26.Mos4.HIV, Protein: subtype C gp140	NCT03060629	HVTN705/Imbokodo	2	Alum	Jul 2022
Vector: Ad26.Mos4.HIV, Protein: bivalent subtype C gp140 and mosaic gp140	NCT03964415	HVTN706/Mosaico	3	Alum	Mar 2024

* Actual or predicted completion date.

Vaccin contre VIH

- « Recombinant canarypox (ALVAC-vCP2438) containing HIV-1 gag (clade B LAI), pro (clade B LAI), env (gp120; clade ZM96.C), and gp41 (clade B LAI) transmembrane anchor », adjuvanté avec MF59
- Essai réalisé en Afrique du Sud
- > 5000 participants
- Proche de l'essai RV144, qui avait montré une efficacité modérée en 2009 en Thaïlande en utilisant « Recombinant canarypox (ALVAC; vCP1521) containing HIV-1 gag (clade B LAI), pro (clade B LAI), env (gp120 AE 92TH023), and gp41 (clade B LAI) transmembrane anchor » adjuvanté aluminium
- Arrêt prématuré

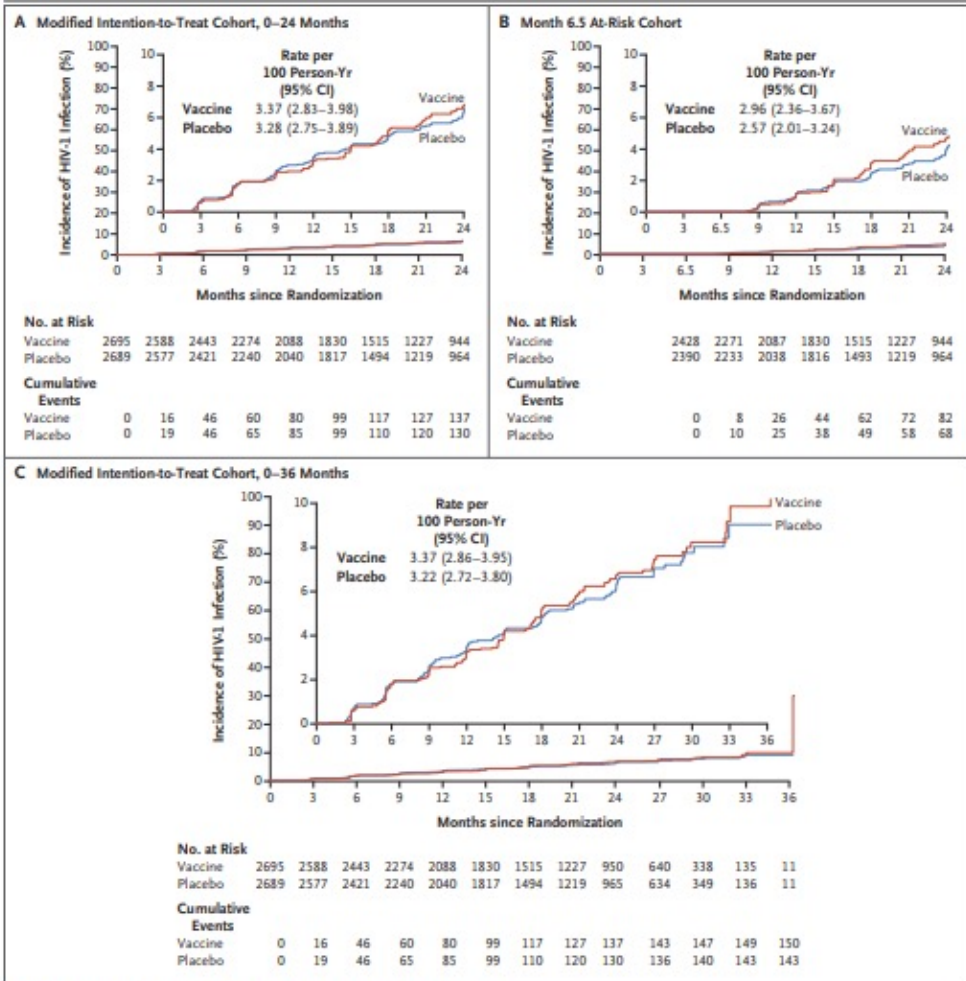


Figure 1. Kaplan–Meier Analysis of HIV-1 Infection in Three Cohorts.

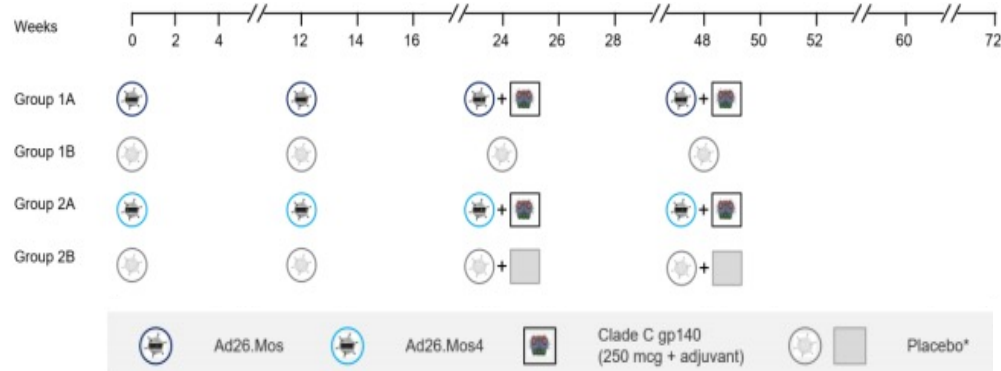
Bekker LG, et al.; N Engl J Med. 2021 Mar 25;384(12):1089-1100.
Zolla Pazner S et al., Lancet HIV 2021; 8: e449–52

Vaccin contre VIH

Safety and immunogenicity of two heterologous HIV vaccine regimens in healthy, HIV-uninfected adults (TRAVERSE): a randomised, parallel-group, placebo-controlled, double-blind, phase 1/2a study

Lancet HIV 2020; 7: e688–98

Lindsey R Baden*, Daniel J Stieh*, Michal Sarnecki, Stephen R Walsh, Georgia D Tomaras, James G Kublin, M Juliana McElrath, Galit Alter, Guido Ferrari, David Montefiori, Philipp Mann, Steven Nijs, Katleen Callewaert, Paul Goepfert, Srilatha Edupuganti, Etienne Karita, Johannes P Langedijk, Frank Wegmann, Lawrence Corey, Maria G Pau, Dan H Barouch, Hanneke Schuitemaker, Frank Tomaka, and the Traverse/HVTN 117/HPX2004 Study Team



- Etude aux USA et Rwanda, 201 patients randomisés, 198 vaccinés
- Vaccin tétravalent contient Ad26-encoded mosaic Env supplémentaire
- Vaccin tétravalent, bon profil de tolérance/ sécurité, idem trivalent
- Tétravalent induit une meilleure réponse immune que le trivalent (Elisa; Ac de liaison, ADCC, IFN gamma, réponse CD4, CD8)
- Poursuite en Phase 3 (NCT03060629 SA, femmes et NCT03964415 USA, Europe MSM)

Vaccin contre le Lyme

- Borréliose de Lyme: 300000 cas annuels aux USA, au moins 100000 en Europe
- « Ancêtres » vaccins ciblant OspA efficaces (1 vaccin recombinant + AI: FDA en 1998 mais trop peu utilisé/polémique, fin commercialisation en 2002) et le 2^{ème} 1 vaccin recombinant sans AI pas de commercialisation
- Plusieurs approches dans les stratégies vaccinales:
 - Cibler *B. burgdoferi* dans le vecteur (OspA)
 - Cibler le spirochète chez l'hôte (protéines de surface, lipides paroi bactérienne, bactéries mutées vivantes)
 - Bloquer la transmission (vaccin ciblant les réservoirs et les vecteurs, générant une immunité anti-tiques)

Gomes-Solecki M, et al., Clin Infect Dis. 2020 :10;70(8):1768-1773.
Steere AC, et al., N Engl J Med. 1998 Jul 23;339(4):209-15.
Sigal LH, et al., N Engl J Med. 1998 Jul 23;339(4):216-22.
Nigrovic LE, et al., Epidemiol Infect. 2007 Jan;135(1):1-8.

Vaccin contre le Lyme

- But= vaccin ciblant les espèces USA et Europe
- Arrêt du vaccin hexavalent Baxter bioscience, NCT01504347,
- Seuls vaccinen phase de développement chez l'Homme: vaccin recombinant hexavalent ciblant 6 sérotypes OspA, vaccin VLA15 recombinant +Aluminium
- Phase 2 Valneva, NCT03769194, NCT03970733
 - Pas de publication complète des résultats
 - Safety ok , taux d'AC élevés
 - Accord avec Pfizer pour la phase 3 pas encore débutée
- Fast track FDA phase 2

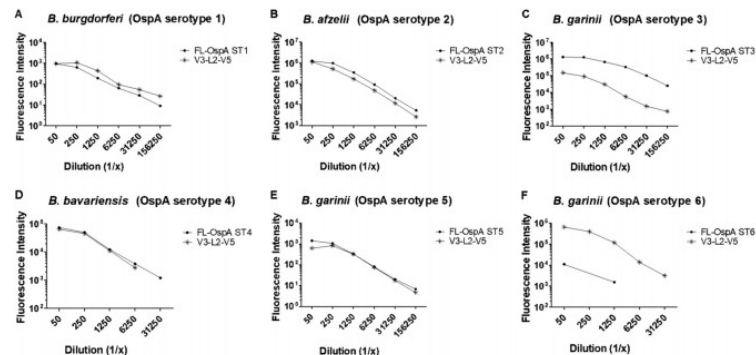


FIG 7 Antibodies generated by V3-L2-V5 versus the corresponding FL-OspA serotypes. The antibodies generated by V3-L2-V5 were tested by surface binding assay. The binding of vaccine-induced antibodies to OspA was compared to the binding of antibodies generated by FL-OspA of the corresponding serotypes (ST1 to ST6). The surface binding assay was carried out with *B. burgdorferi* OspA ST1 Z57, *B. afzelii* OspA ST2 Pra10, *B. garinii* OspA ST3 PFr, *B. bavariensis* OspA ST4 PFin, *B. garinii* OspA ST5 PHei, and OspA ST6 KL11. The results are represented as fluorescence intensity.

Wressnigg N, et al., Clin Vaccine Immunol. 2014 Nov;21(11):1490-9

Nayak A, et al., Infect Immun. 2020 Mar 23;88(4):e00917-19

<https://clinicaltrials.gov/ct2/show/results/NCT03769194?term=VLA15&cond=Lyme&draw=2&rank=3>

Vaccin contre le VRS

- Vrai problème chez le nourrisson: 3,2 millions d'hospitalisations dans le monde en 2015, 118000 décès, 50% < 6mois
- Femmes enceintes
- Adultes: personnes âgées, immunodéprimés: formes graves, 4-10% mortalité
- Histoire vaccinale compliquée

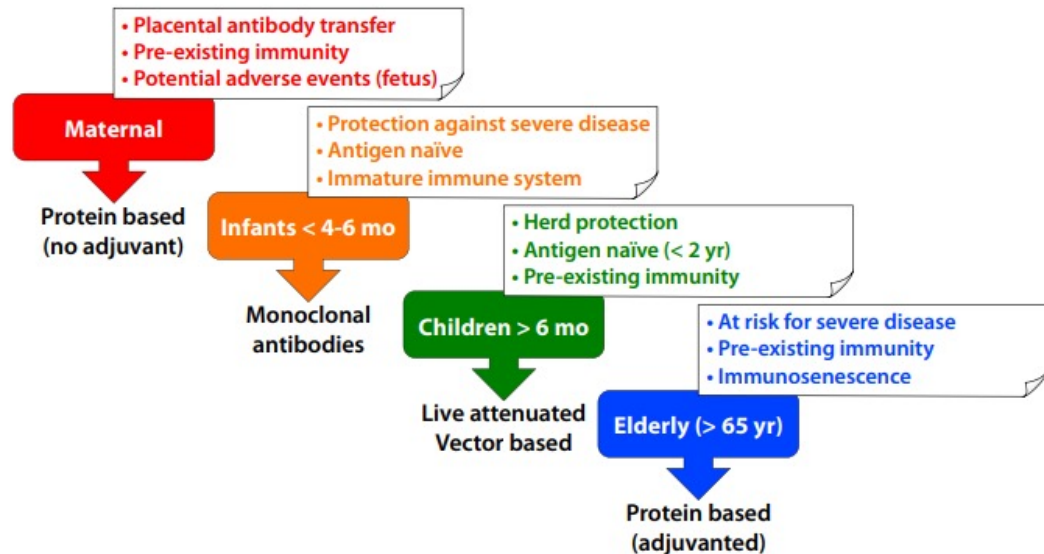


Figure 3. Target populations and respiratory syncytial virus vaccine types. There are different vaccine strategies according to the main target populations. The 4 main target populations are color coded. The nuances and characteristics of each target population are included in the adjacent balloon and the preferred vaccine strategy underneath each target population.

Vaccin contre le VRS

RSV Vaccines in Clinical Development

Vaccine type (manufacturer)	Viral target	Target population	Administration route	Clinical development	Advantages	Challenges
PROTEIN VACCINES						
Particle based						
RSV F nanoparticle (Novavax, Gaithersburg, Maryland)	Prefusogenic	Maternal, elderly, pediatric	Systemic	Phase 3, phase 2, phase 1	Safe, immunogenic	Post-F based? Risk of ERD, antibody durability
Subunit						
DS-Cav1 (NIH/NIAID, Bethesda, Maryland)	Pre-F	Maternal and elderly	Systemic	Phase 1	Induce high-affinity neutralizing antibody, facilitate cross-priming, safe	Factors that affect transplacental transfer, instability of pre-F, antibody durability, no protection for premature infants
GSK RSV F (GlaxoSmithKline, Brentford, United Kingdom)	Pre-F	Maternal and elderly	Systemic	Phase 1		
DPX-RSV (Immunovaccine, Dartmouth, Canada, and VIB, Flanders, Belgium)	SH	Elderly	Systemic	Phase 1		
RSV-F (Janssen, Beerse, Belgium)	Pre-F	Elderly	Systemic	Phase 1		
RSV-F (Pfizer, New York, New York)	Pre-F	Maternal and elderly	Systemic	Phase 2		
RSV-G (Advaccine Biotech, Beijing, China)	G	Pediatric and elderly	Systemic	Phase 1		
LIVE VACCINES						
Vector based						
AdV26 RSV (Janssen)	Pre-F	Pediatric and elderly	Systemic	Phase 2	Not attenuated, low risk of ERD, no interference with maternal antibodies	Potential for developing antivector immunity
ChAdV155-RSV (GlaxoSmithKline)	Pre-F, N, M2-1	Pediatric	Systemic	Phase 2		
VXA-RSV (AdV5) (Vaxart, South San Francisco, California)	Post-F	Elderly	Mucosal and systemic	Phase 1		
MVA-BN RSV (Bavarian Nordic, Kvistgaard, Denmark)	Post-F, GA/GB, N, M2	Elderly	Systemic	Phase 2		
Live-attenuated/chimeric						
rBCG/N-hRSV (Universidad de Chile, Santiago, Chile)	N	Newborn	Systemic	Phase 1	Predominant T _H 1 immune responses	
RSV/ΔG (Intravac)	Lacks G	Pediatric	Mucosal	Phase 1	Low risk of ERD, intranasal delivery, replication in presence of maternal antibody, broad stimulation of immune responses	Balance of attenuation/immunogenicity, reverse to wild type, stability for mass production
RSV ΔNS2 Δ1313/1314L RSV 276	Pre-F/post-F	Pediatric	Mucosal and systemic	Phase 1		
RSV 6120/ΔNS2/1030 _s (Sanofi Pasteur, Lyon, France, and NIH)						
SeV/RSV (St Jude Hospital, Atlanta, Georgia)	F	Pediatric	Mucosal	Phase 1		

Abbreviations: Adv, adenovirus; ERD, enhanced RSV disease; F, fusion; G, attachment; MVA, modified vaccinia Ankara virus; ND, not disclosed; NIAID, National Institute of Allergy and Infectious Diseases; NIH, National Institutes of Health; post-F, postfusion; pre-F, prefusion; RSV, respiratory syncytial virus; SeV, Sendai virus; SH, small hydrophobic.

Mejias A, et al., Ann Allergy Asthma Immunol. 2020 Jul;125(1):36-46.
Madhi SA, et al., N Engl J Med. 2020 Jul 30;383(5):426-439.

Vaccin contre *E. coli* uropathogènes

TABLE 1 | Non-antibiotic therapeutic options for the treatment of urinary tract infections.

Therapeutic options	References	Mechanism	Benefits	Drawbacks
Vaccine				
Targeting adhesion	(O'Hanley et al., 1985; De Ree and Van den Bosch, 1987; Riegman et al., 1988; Wizemann et al., 1999; Langermann et al., 2000; Roberts et al., 2004; Poggio et al., 2006; Habibi et al., 2016)	<ul style="list-style-type: none"> • Block the liaison adhesin-host cell receptor (pili vaccine) • Reduction of adhesion and protection against cystitis (FimH vaccine) 	<ul style="list-style-type: none"> • Decrease the bacterial colonization • Protection of the bladder and the kidneys 	<ul style="list-style-type: none"> • Heterogeneity of the proteins of the bacterial membrane
Targeting capsule	(Kajiser et al., 1983; Roberts et al., 1993; Kumar et al., 2005; Stenutz et al., 2006)		<ul style="list-style-type: none"> • Promising animal model results 	<ul style="list-style-type: none"> • No human studies • Great heterogeneity in antigen used making creation of a vaccine with broad protection difficult
Targeting toxins	(O'Hanley et al., 1991; Ellis and Kuehn, 2010)	<ul style="list-style-type: none"> • Reduction of renal injury 	<ul style="list-style-type: none"> • Decrease virulence 	<ul style="list-style-type: none"> • No long-term protection
Targeting iron metabolism	(Alteri et al., 2009; Brumbaugh et al., 2013)	<ul style="list-style-type: none"> • Effective immunologic reaction against specific molecules 	<ul style="list-style-type: none"> • Protection of the bladder and the kidneys • Reduce UTI recurrence 	<ul style="list-style-type: none"> • Cannot target all UPEC strains (heterogeneity of the targets)

Vaccin contre *E. coli* uropathogènes

Table 1 – Available vaccines, administration methods, and vaccine content

Vaccine	Method of administration	Bacterial content
UroVaxom (OM-89)	One oral tablet to be taken once a day for 3 mo ± booster tablet for the first 10 d of months 6–9	6 mg of lyophilised bacterial lysates derived from 18 <i>E. coli</i> strains
Uromune	Two doses of 100 µl each (10 ⁸ bacteria/puff) daily sublingually, for a duration of 3 mo	<i>E. coli</i> , <i>Klebsiella pneumoniae</i> , <i>Proteus vulgaris</i> , <i>Enterococcus faecalis</i>
Solco-Urovac	Vaginal suppository given weekly for the first 3 wk, then a booster monthly for 3 mo Intramuscular injection, initially weekly for 3 wk, with a booster at 6 mo	10 Uropathogenic strains of bacteria including 6 <i>E. coli</i> strains, <i>K. pneumoniae</i> , <i>Proteus mirabilis</i> , <i>Proteus morganii</i> , and <i>E. faecalis</i>
ExPEC4V	Single intramuscular injection of 0.5 ml	Genetically detoxified form of exotoxin A from <i>Pseudomonas aeruginosa</i> linked to four serotype surface polysaccharide antigens of <i>E. coli</i> (O1A, O2, O6A, O25B)

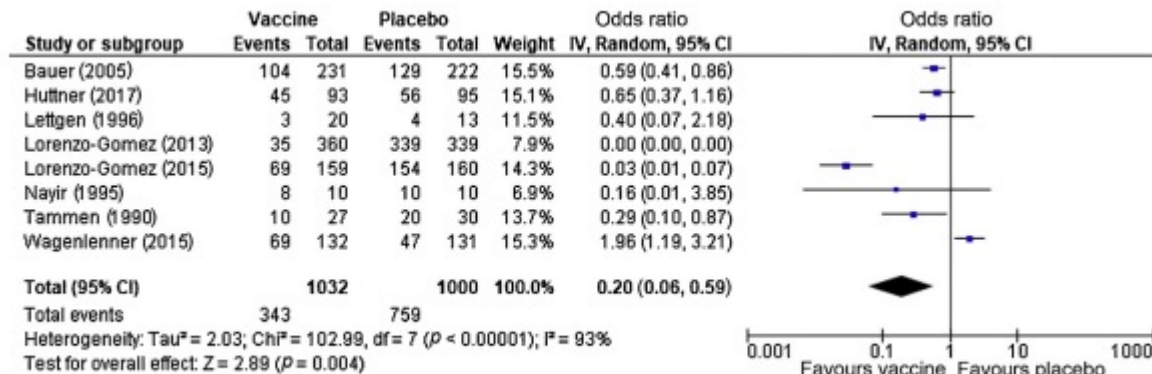


Fig. 5 – Long-term efficacy of vaccines. CI = confidence interval; IV = inverse variance.

Vaccin contre *Staphylococcus aureus*



Staphylococcus aureus Vaccine Research and Development: The Past, Present and Future, Including Novel Therapeutic Strategies

Jonah Clegg^{1,2}, Elisabetta Soldaini¹, Rachel M. McLoughlin², Stephen Rittenhouse³, Fabio Bagnoli¹ and Sanjay Phogat^{1*}

TABLE 1 | *Staphylococcus aureus* vaccines currently enrolled in clinical trials.

Company	Vaccine	Phase	Clinical trial number	Study population	Literature
GSK	SA-5Ag: Adjuvanted	I: Recruiting	NCT04420221	18 – 50 year olds at risk of recurrent skin infections	
Novadigm Therapeutics	NDV-3A: Als-3 (<i>C. albicans</i> cross reactive cell wall protein) + Alum	II: Ongoing	NCT03455309	Military Personnel	(136, 137)
Olymvox	rFSAV: Hla, SpA, SEB, IsdB, MntC + Alum	II: Ongoing	CTR20181788, NCT03966040		(138)
Pfizer	SA4Ag: CP5-dptx, CP8-dptx, CifA, MntC	IIb: Failure	NCT02388165	Patients undergoing spinal surgery	(20, 139–141)
Integrated Biotherapeutics	i. Stebvax: SEB + alum ii. IBT-V02: SEB, SEA, TSST-1, LukS, LukF, LukAB, Hla + alum	I: Completed I: Scheduled	NCT00974935	18 – 40 year olds	(142)

Le plus attendu.....



Le vaccin contre la connerie: toujours pas au point
<http://ministere Duchomage.fr>



<https://nostalgia.blog4ever.com>

