

## Mountebanks...

### • Old Oxford Dictionary:

- A charlatan, a person who falsely claims knowledge of or skill in some matter, esp. for personal gain; a person who pretends to be something he or she is not, in order to gain prestige, fame, etc.
- An itinerant quack who from an elevated platform appealed to his audience by means of stories, tricks, juggling, and the like, in which he was often assisted by a professional clown or fool.
- Italian: monta inbanco
- Montare = to ascend or go up to a place,
- Banco = a bench ["To mount on a bench"]

https://www.oed.com/dictionary/mountebank\_n (accessed 12/4/25) https://hillcantons.blogspot.com/2010/10/mountebank-class-take-two.html (accessed 12/4/25) html (accessed 12/



# Toady / Toadies...

- · Old Oxford Dictionary:
  - One who eats toads; originally the attendant of a charlatan, employed to eat or pretend to eat toads (held to be poisonous) to enable the charlatan to exhibit skill in expelling poison
  - figurative. A fawning flatterer, parasite, sycophant



https://www.oed.com/dictionary/toad-eater\_n (accessed 12/4/25)
https://pulterproject.northwestern.edu/curations/c72-what-is-a-mountebank.html (accessed 12/4/25)
https://www.amazon.com/Posteraza:-Mountebank-Engraving-English-Century/slp/807C30ZMHM (accessed 12/4/25)





## **Disclosures**

- Dr. Aeschlimann has no relevant financial relationships to disclose
- This activity may contain discussion of unlabeled/unapproved use of drugs.
  - The content and views presented in this educational program are those of the faculty and do not necessarily represent those of the University of Connecticut School of Pharmacy.
  - Please refer to the official prescribing information for each product for discussion of approved indications, contraindications, and warnings

# **Disclosures, Part 2**

- The use of public-domain materials found on the CDC and its Advisory Committee on Immunization Practices (ACIP) websites does not imply endorsement by CDC, ACIP, ATSDR, HHS or the United States Government of Dr. Aeschlimann, UConn, and/or UConn School of Pharmacy
  - Similarly, Dr. Aeschlimann's use of materials from those websites does not imply his blind endorsement of the policies and actions of those Government agencies, nor does it mean that the information on those websites is considered "evidence-based" or "scientifically correct"
- Additionally, any reference to specific commercial products, manufacturers, companies, or trademarks does not constitute its endorsement or recommendation by the U.S. Government, Department of Health and Human Services, or Centers for Disease Control and Prevention
- The public-domain materials presented have not been substantively changed, and all of the source materials are available on the agency's website for no charge.

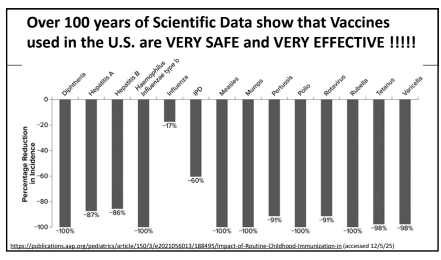
# **Learning Objectives**

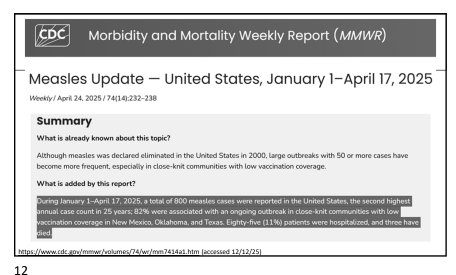
- At the conclusion of this CPE activity, participants should be able to:
  - 1) Describe at least one important change (or proposed change) in childhood and adult vaccination recommendations put forth by the CDC and/or ACIP
  - Given a patient who inquires about receiving respiratory virus or bacteria vaccinations (e.g., Influenza, COVID-19, Respiratory Syncytial Virus (RSV), Pneumococcal), outline important differences between multiple products when they exist
  - Identify evidence-based pharmacotherapeutic treatments for common vaccine-preventable illnesses

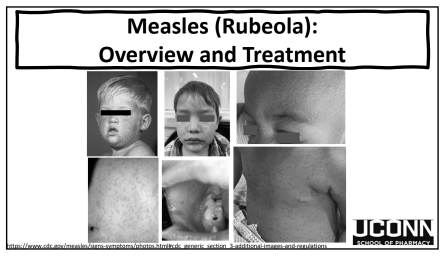
# **Pre-Test Time!**

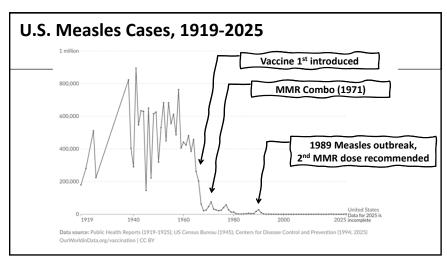
10

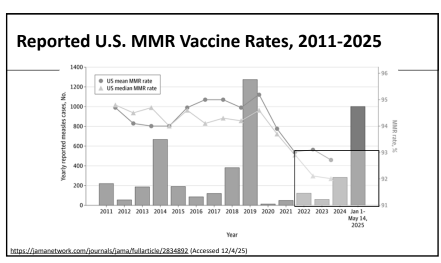
- The U.S. Dept. of HHS, the CDC, the FDA, and the Advisory Committee on Immunization Practices (ACIP) have been infiltrated by Mountebanks, Antivax Cranks, Grifters, Charlatans, & Toadies...
- Which **ONE** of the following is **NOT** a recommendation that has been **APPROVED** by **ACIP** in **2025**?
  - a) All adults receive seasonal influenza vaccines only in single dose formulations that are free of thimerosal as a preservative
  - b) State and local jurisdictions should require a prescription for the administration of a COVID-19 vaccination
  - c) COVID-19 Vaccination should be based on individual-based decision-making (Shared Clinical Decision Making) for Adults 65 and older
  - d) Universal Hepatitis B Vaccination at birth is no longer recommended for all newborns
  - e) For Hepatitis B vaccination, post-vaccine serology results should determine need for subsequent doses (in the 3-dose series)

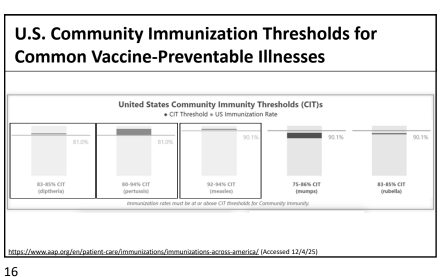


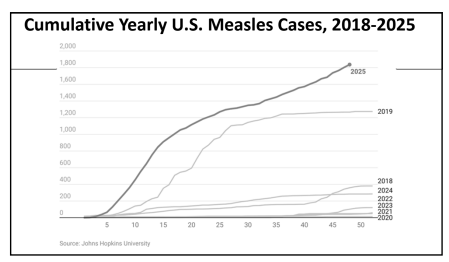


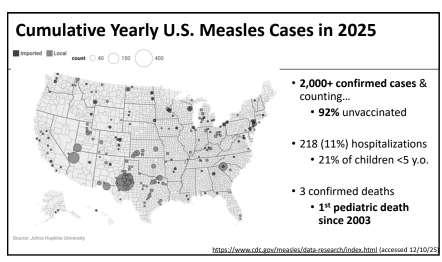


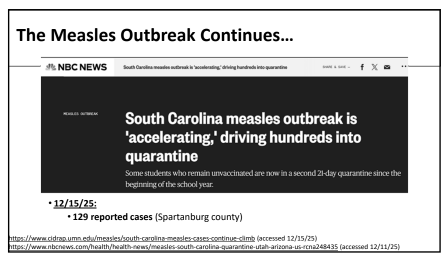


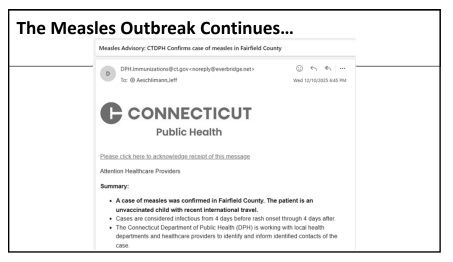












# **Measles: Key Facts**

- HIGHLY CONTAGIOUS (~90% infection following exposure if susceptible)
  - Airborne, person-to-person contact
- Infection course: Prodrome phase → Exanthem phase → Recovery/Immunity
- Contagious <u>+</u> 5 days relative to rash appearance
- Complications (30% of cases):
  - Immunosuppression / Secondary infections
  - Diarrhea
  - Pneumonia
  - Encephalitis, Acute disseminated encephalomyelitis (ADEM)

# **Treatment of Measles**

- No approved antiviral therapies
  - Ribavirin ???
- Supportive care:
  - Antipyretics
  - Fluids/Nutrition
  - Monitoring, diagnosis, & treatment of 2° infections
- Vitamin A therapy ???
- Chin-Ups at Airports???

22



## **Measles Treatment: Vitamin A?**

- Serum Vitamin A levels ↓ during measles infection
- Pre-existing deficiency + malnutrition w/ acute measles infection:
  - Possible delayed recovery, † pneumonia mortality
  - Xerophthalmia → Corneal dryness, conjunctival keratinization
- Barclay, et al. (BMJ 1987;294(6567):294.)
  - 180 hospitalized children in rural Tanzania
  - 90% with low serum Vit. A
  - Vit. A (200,000 IU) orally x 2 doses
  - ↓ mortality in < 2 y.o. (1/46 vs. 7/42)

(months)	vitamin A
<9	14
9-11	12

TABLE I-Mortality of children admitted with measles

Age (months)	Given vitamin A	Controls	Given vitamin A	Controls
<9	14	9		2 (22)
<9 9-11 12-23	12	10	1 (5)	2 (20)
24.35	11	16	3 (27)	2(13)
36-47 48-59 ≽60	11	13 6	1 (9) 1 (13)	1 (8)
≥60	12	15	1(15)	2(13)
Total	. 88	92	6 (7)	12 (13)

24

https://pubmed.ncbi.nlm.nih.gov/3101849/

**Measles Treatment: Vitamin A?** Table 3. Mortality and Morbidity in 189 Children with Measles, According to Treatment Group.\* - A Randomized, Controlled Trial of Vitamin A in RELATIVE RISK (95% CI)† Children with Severe Measles P VALUE 0.046 0.21 (0.05-0.94) 10 Age at death Randomized, double-blind trial in South Africa • 189 children < 13 y.o. hospitalized with acute Pneumonia (days)
Duration
≥10 measles complicated by pneumonia, diarrhea, Diarrhea (days)
Duration
≥10 Vit. A (200,000 I.U.) orally x 2 doses 0.51 (0.28-0.92) 0.23 (0.05-1.06) Herpes stomatitis 0.38 (0.13-1.16) Intensive care · Results: Adverse outcome 15.24 (8, 11, 19) 10.52 (7, 9, 13) • ~80% were < 2 y.o, ~66% < 1 y.o. \*In the columns representing the treatment groups, the values in italics are means, followed in par ercentiles, medians, and 75th percentiles. All other values are numbers of patients. The concentration medians, and 75th percentiles. All other values are numbers of patients.

\*Relative risk denotes the ratio of the incidence of an event in the vitamin A group to the incidence of the event in the placebo group. Cl denotes confidence interval.

\*\*In Aura in Auraion.\*\* postmessles croup, or transfer for Serum Vit. A ↓ lower limit of normal in

## **Measles Treatment: Vitamin A?**

- Cochrane Systematic Review (2005):
  - "To determine whether vitamin A, commenced after measles has been diagnosed, prevents mortality, pneumonia or other complications in children."
- 8 studies (2574 participants) from 1932-1999, 6 blinded
  - Africa (n=6), Japan (n=1), England (n=1)
  - Generally heterogeneous studies

https://pmc.ncbi.nlm.nih.gov/articles/PMC7076287/

## **Measles Treatment: Vitamin A?**

- Cochrane Systematic Review (2005):
  - Pooled results from 7 high-quality studies:
    - No significant effect on mortality (RR 0.83, CI 0.51-1.34)
  - Results from 3 high-quality studies in hospitalized children in high case-fatality areas:
    - Significant 64% ↓ in mortality (RR 0.40, CI 0.19-0.87)
    - Driven by 83% ↓ mortality in children < 2 y.o (RR 0.21, CI 0.07-0.66)

https://pmc.ncbi.nlm.nih.gov/articles/PMC7076287/

# Measles Treatment: CDC Recommendations

- · Vitamin A:
  - "...does not prevent measles and is not a substitute for vaccination."
  - "...may be administered to infants and children in the United States with measles under the supervision of a healthcare provider as part of supportive management."
  - "...it should be administered immediately upon diagnosis and repeated the next day for a total of 2 doses."
    - 50,000 IU for infants younger than 6 months of age
    - 100,000 IU for infants 6–11 months of age
    - 200,000 IU for children 12 months of age and older

 $\underline{https://www.cdc.gov/measles/media/pdfs/2025/05/hcp-caring-for-patients-measles-fact-sheet.pdf}$ 

# Measles Treatment: CDC Recommendations

• Ribavirin:

28

- "...demonstrates in vitro activity against measles virus."
- "While ribavirin has been used to treat patients with severe measles disease or severely immunocompromising conditions, clinical data are lacking regarding its efficacy."
- "...is not approved by the U.S. Food and Drug Administration (FDA) to treat measles."

 $\underline{https://www.cdc.gov/measles/media/pdfs/2025/05/hcp-caring-for-patients-measles-fact-sheet.pdf}$ 

# Measles Treatment: CDC Recommendations

### • Antibiotics:

- "There is no evidence to support routine use of antibiotics for measles treatment."
- "Measles may be complicated by secondary bacterial infections for which antibiotic treatment is indicated."

 $\underline{https://www.cdc.gov/measles/media/pdfs/2025/05/hcp-caring-for-patients-measles-fact-sheet.pdf}$ 

# Measles Treatment: CDC Recommendations

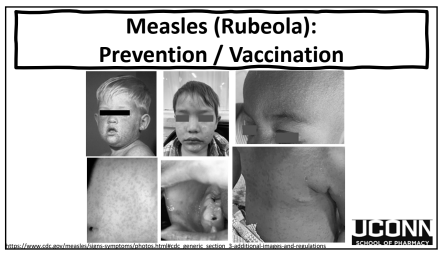
### Isolation:

• "Infected people should be isolated for 4 days after they develop a rash; airborne precautions should be followed in healthcare settings."

### Vaccination:

- "MMR vaccination is the best way to prevent measles and its complications."
- "People exposed to measles may be eligible for post-exposure prophylaxis with MMR vaccine within 72 hours (or immunoglobulin within 6 days)"

 $\underline{\text{https://www.cdc.gov/measles/media/pdfs/2025/05/hcp-caring-for-patients-measles-fact-sheet.pdf}$ 



Recommende	d Child ar	d Adoles	scent In	nmuniza	ation Sched	ule	Amorican	A andomy	of Dodintu	ine Co
Table 1 for Ages 18 Ye	ars or You	nger, Un	ited Sta	tes, 202	.5		American DEDICATED TO	,		(Marile)
These recommendations must be read with the opportunity as indicated by the outlined purple l										
Vaccine and other immunizing agents	Birth 1 m	os 2 mos 4	mos 6 mo	s 8 mos 9 n	nos 12 mos 15 m	nos 18 mos 19-	-23 mos 2-3 yrs 4-6	yrs 7–10 yrs 11–	12 yrs   13–15 yrs	16 yrs   17–18 yrs
Respiratory syncytial virus (RSV-mAb [nirsevimab, clesrovimab])	1 dose during RSV vac	RSV season dependi cination status (See	ng on maternal • Notes)	1 dose nir	sevimab during RSV seas	on (See Notes)				
Hepatitis B (HepB)	1º dose	2 <sup>rd</sup> dose			3° dose					
Rotavirus (RV): RV1 (2-dose series), RV5 (3-dose series)		1" dose	2 <sup>nl</sup> See dose Notes							
Diphtheria, tetanus, and acellular pertussis (DTaP <7 yrs)		1" dose	2" 3" dose			4º dose	5.4	ose		
Haemophilus influenzae type b (Hib)		1" dose	2 <sup>nd</sup> See dose Notes		3 <sup>rd</sup> or 4 <sup>th</sup> dose (See Notes)					
Pneumococcal conjugate (PCV15, PCV20)		1" dose	2" 3" dose dose		4º dose					
Inactivated poliovirus (IPV)		1" dose	2 <sup>rd</sup> dose		3° dose		4.6	254		See
COVID-19 (1vCOV-mRNA, 1vCOV-aPS)				1 or more	doses of 2025-2026 vac	cine (See Notes)	GRRRRR	1 dose of 2025-2026		
Influenza					1 or 2 dose	es annually (See Notes)			1 dose annually (\$	ee Notes)
Measles, mumps, and rubella (MMR)				See Notes	1" dose		2"6	**		
Varicella (VAR)					1" dose		2"6	954		

## **Measles Prevention: Vaccine Products**

- MMR [Measles-Mumps-Rubella]:
  - M-M-R II®
  - PRIORIX®
- MMRV [Measles-Mumps-Rubella-Varicella]:
  - ProQuad®
- ALL are FDA-approved for use in children 12 months 12 years of age

# Measles Prevention: CDC Recommendations (before Oct. 2025)

Routine Vaccination Recommendations to Protect against Measles

#### Children

34

CDC recommends two doses of measles-containing vaccine routinely for children, starting with the first dose at age 12 through 15 months and the second dose at age 4 through 6 years before school entry. This can be administered as MMR or MMRV vaccine. Children can receive the second dose of MMR vaccine earlier than 4 through 6 years, as long as it is at least 28 days after the first dose. A second dose of MMRV vaccine can be given 3 months after the first dose up to 12 years of age.

CDC recommends that separate <u>MMR</u> and varicella vaccines be given for the first dose in children aged 12–47 months; however, MMRV may be used if parents or caregivers express a preference.

https://www.cdc.gov/vaccines/vpd/mmr/hcp/recommendations.html (accessed 12/4/25)

## **Measles Prevention – ACIP Changes its Recommendations to CDC for Routine Vaccination**

#### 9/18/2025 ACIP Meeting:

Voted (8-for, 3-against, 1-abstain) to REMOVE the option to use MMRV in children 12-47

Vote: MMRV Vaccines Vote
Dr. Martin Kullidorfi (ACIP Chair) read the following proposed ACIP voting language for MMRV vaccines into the record:

v accures more rescore. The pediatric vaccine schedule should be updated to reflect the following change: -For measles, mumps, rubella and varicella vaccines given before age 4 years, the combined MMRV vaccine is not recommended. -Children in this age group should receive separate measles, mumps, and rubella vaccine and variental vaccine (MMRVV).

Hillary Blackburn, PharmD, M.B.A.

Director of Medication Access and Affordability, AscensionRx, Term: 9/11/2025-

Hillary Blackburn, PharmD, M.B.A., leads initiatives to optimize medication access for underserved populations and improve affordability in value-based care. She previously served as Chief Pharmacy Officer at the Dispensary of Hope, overseeing formulary and research strategy. She is also a leader in professional pharmacy organizations, host of the Talk to Your Pharmacist podcast, and author of How Pharmacists Lead.

Hilary Blackburn, a pharmacist based in St. Louis, Missouri. She is the daughter-in-law of Republican Tennessee Senator Marsha Blackburn.

35

#### Motion/Vote: MMRV Vaccines

varicella vaccine (MMR+V)."

Dr. Pagano seconded the motion. No COIs were declared. The motion carried with 8 votes in favor, 3 votes opposed, and 1 abstention. The disposition of the vote was as follows:

8 Favored: Pagano, Mihoan, Stein, Griffin, Pollak, Pebsworth, Levi, Kulldorff 3 Opposed: Blackburn, Hibbeln, Meissner 1 Abstained: Malone

https://www.cdc.gov/acip/downloads/minutes/summary-2025-9-18-19-508.pdf (Accessed 12/5/25

## **Measles Prevention – ACIP Changes its Recommendations to CDC for Routine Vaccination**

- 9/18/2025 ACIP Meeting:
  - Voted (8-for, 3-against, 1-abstain) to **REMOVE** the option to use MMRV in children 12-47 months old
- Why?????

36

- Original CDC recommendations (from 2009 ACIP) based off slightly higher risks of fevers/febrile seizures for MMRV vs. separately-administered doses of MMR & VARIVAX [evaluated at 5 & 12 days]...
- ~1 extra febrile seizure for every 2,300-2,600 administered MMRV doses
- · No significant new scientific data presented at Sept. 2025 ACIP meeting...

 $\underline{https://www.cidrap.umn.edu/childhood-vaccines/new-cdc-advisers-scale-back-recommendations-mmrv-vaccine-young-kids} \ (accessed 12/5/25) \ (accessed 12/5$ 

https://www.cdc.gov/vaccines/vpd/mmr/hcp/vacopt-faqs-hcp.html (accessed 12/5/25)

# ProQuad® Fever & Febrile Seizures

### • <u>Pre-Marketing</u> <u>Comparative Data:</u>

 Two children (of ~2,700) had febrile seizures after ProQuad dose #2

 These children appeared to have concurrent viral illness during vaccination Table 1: Vaccine-Related Injection-Site and Systemic Adverse Reactions
Reported in ≥1% of Children Who Received ProQuad Dose 1 or M-M-R II and VARIVAX
at 12 to 23 Months of Age (0 to 42 Days Postvaccination)

Adverse Reactions	ProQuad (N=4497) (n=4424) %	M-M-R II and VARIVAX (N=2038) (n=1997) %
Injection Site*		
Pain/tenderness/soreness <sup>†</sup>	22.0	26.7
Erythema <sup>†</sup>	14.4	15.8
Swelling <sup>†</sup>	8.4	9.8
Ecchymosis	1.5	2.3
Rash	2.3	1.5
Systemic		
Fever <sup>†,‡</sup>	21.5	14.9
Irritability	6.7	6.7
Measles-like rash <sup>†</sup>	3.0	2.1
Varicella-like rash <sup>†</sup>	2.1	2.2
Rash (not otherwise specified)	1.6	1.4
Upper respiratory infection	1.3	1.1
Viral exanthema	1.2	1.1
Diarrhea	1.2	1.3

- \* Injection-site adverse reactions for M-M-R II and VARIVAX are based on occurrence with either of the vaccines administered.
- Designates a solicited adverse reaction. Injection-site adverse reactions were solicited only from Days 0 to 4 postvaccination.
- <sup>‡</sup> Temperature reported as elevated (≥102°F, oral equivalent) or abnormal.
- N = number of subjects vaccinated.
  n = number of subjects with safety follow-up.

https://www.merck.com/product/usa/pi\_circulars/p/proquad/proquad\_pi.pdf

### **ProQuad® Fever & Febrile Seizures**

- Post-Marketing Observational Safety Surveillance Study:
  - Age-, gender-, and date-of-vaccination- (day and month) matched subjects
  - Given M-M-R II and VARIVAX concomitantly
- Differences only observed following Dose #1 within the 5-12d evaluation range:

Table 11: Confirmed Febrile Seizures Days 5 to 12 and 0 to 30 After Vaccination with ProQuad (dose 1) Compared to Concomitant Vaccination with M-M-R II and VARIVAX (dose 1) in Children 12 to 60 Months of Age

Time Period	ProQuad cohort (N=31,298)			MR+V cohort N=31,298)	Relative risk (95% CI)
	n	Incidence per 1000	n	Incidence per 1000	
5 to 12 Days	22	0.70	10	0.32	2.20 (1.04, 4.65)
0 to 30 Days	44	1.41	40	1.28	1.10 (0.72, 1.69)

https://www.merck.com/product/usa/pi\_circulars/p/proquad/proquad\_pi.pdf

38

# Measles Prevention – ACIP Changes its Recommendations to CDC for Routine Vaccination

- What does this mean practically???
  - CDC adopted the ACIP recommendations on 10/6/25
  - Two separate injections now required for the first MMR / Varicella vaccinations...
  - MMRV may not be covered for children under 4 y.o. in the Vaccines for Children program...
    - But...only ~15% of parents opt to use MMRV for the 12-15 month dose
    - State Medicaid programs & Private Insurers may still choose to cover MMRV

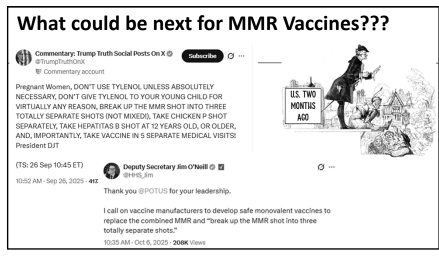
https://www.cidrap.umn.edu/childhood-vaccines/new-cdc-advisers-scale-back-recommendations-mmrv-vaccine-young-kids (accessed 12/5/25) https://www.astho.org/communications/blog/2025/downstream-effects-of-cdc-adopting-acip-recommendations/ (accessed 12/5/25)

## **Measles Prevention: Outbreak Situations**

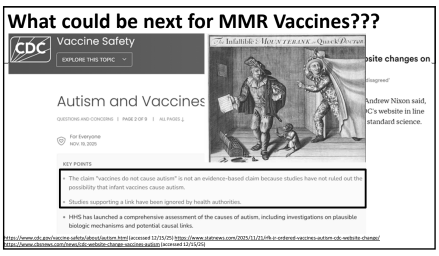
- 2022 ACIP Recommendations:
  - · General recommendation:
    - Infants aged 6-11 months should receive a single dose of MMR
  - Post-Exposure prophylaxis ("PEP"):
    - Unvaccinated persons should receive 1 dose of MMR within 72 hours of exposure to a person with infectious measles
    - Complete the 2-dose MMR series ≥ 28 days later
  - · Product labelling "Fascinoma":
    - M-M-R II use in measles PEP is "ON-label"
    - PRIORIX® use in measles PEP is "OFF-label"

https://www.cdc.gov/mmwr/volumes/71/wr/mm7146a1.htm

40







December 5, 2025: Changes in ACIP Hepatitis B Vaccination Recommendations



# 12/5/25 ACIP Meeting Addressing Universal Hepatitis B Vaccine Birth-Dose for Infants

- Vote #1:
  - · For infants born to HBsAg-negative women:
    - ACIP recommends individual-based decision-making, in consultation with a health care provider, for parents deciding when or if to give the HBV vaccine, including the birth dose. (1) For those not receiving the HBV birth dose, it is suggested that the initial dose is administered no earlier than 2 months of age. Y/N
      - (1) Parents and health care providers should consider vaccine benefits, vaccine risks, and infection risks. Parents and health care providers should also consider whether there are risks, for example, such as a household member is HBsAg-positive or when there is frequent contact with persons who have emigrated from areas where Hepatitis B is common.

https://www.cnn.com/health/live-news/cdc-vaccine-meeting-hepatitis-b-12-05-25 (accessed 12/5/25)

## Results of the 12/5/25 ACIP Meeting & Vote Details

Vote #2:

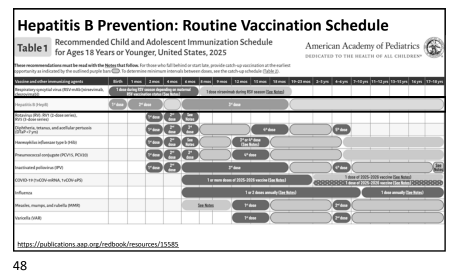
46

 When evaluating the need for a subsequent HBV vaccine dose in children, parents should consult with health care providers to determine if a post-vaccination anti-HBs serology testing should be offered. Serology results should determine whether the established protective anti-HBs titer threshold of ≥10 mIU/mL has been achieved. The cost of this testing should be covered by insurance. Y/N

 $\underline{https://www.cnn.com/health/live-news/cdc-vaccine-meeting-hepatitis-b-12-05-25} \ (accessed \ 12/5/25)$ 

# **Hepatitis B Infection: Key Facts**

- Incurable viral infection
- Transmitted through blood & various body fluids
- High risk for perinatal transmission (up to 90%)
- Can survive on surfaces for weeks
- ~90% of infants infected with HepB will develop chronic infection
- Chronic infection leads to:
  - Liver cirrhosis, hepatocellular carcinoma, death



# **Hepatitis B Prevention:** Vaccine Products

- Recombivax HB (1986)
- Engerix-B (1989)
- Protective antibody responses [anti-Hbs > 10 mIU/mL]:
  - ~25% after 1st dose
  - ~63% after 2<sup>nd</sup> dose
  - ~95% after 3<sup>rd</sup> (final) dose

## **Perinatal Hepatitis B Prevention: History and Progress**

ACIP hepatitis B screening recommendations for pregnant women and perinatal postexposure recommendations for infants, United States

Recommendation	Current?	Date of Initial Recommendation	Date Recommendation Superseded or Modified
Perinatal strategies			
Screening and testing for pregnant women			
Universal HBsAg screening in first trimester	Yes	1988 <sup>1</sup>	
Test for HBsAg later in pregnancy for risk behaviors or acute hepatitis	Yes	1988 <sup>1</sup>	
Test for HBsAg at delivery if status is unknown	Yes	1988¹	

• [HBsAg = Hepatitis B Surface Antigen]

https://iournals.sagepub.com/doi/10.1177/00333549231175548 (accessed 12/5/25) https://www.cdc.gov/acip/downloads/slides-2025-09-18-19/02-langer-hep-b-508.pdf (accessed 12/5/25)

49

# **Perinatal Hepatitis B Prevention: History and Progress**

Recommendation Perinatal strategies	Current?	Date of Initial Recommendation	Date Recommendation Superseded or Modified
Post-exposure prophylaxis for infants born to HBsAg (+) pregnant women			
Administer HBIG within 12 hours of birth and HepB vaccine simultaneously or within 7 days of birth	No	1984²	1987³
Administer HBIG and HepB vaccine at birth*	No	1987³	1988¹
Administer HBIG and HepB vaccine within 12 hours of birth*	Yes	1988 <sup>1</sup>	

https://journals.sagepub.com/doi/10.1177/00333549231175548 (accessed 12/5/25) https://www.cdc.gov/acip/downloads/slides-2025-09-18-19/02-langer-hep-b-508.pdf (accessed 12/5/25)

# Perinatal Hepatitis B Prevention: History and Progress

Recommendation Perinatal strategies	Current?	Date of Initial Recommendation	Date Recommendation Superseded or Modified
Infants born to HBsAg status unknown pregnant women			
In populations where screening is not feasible, administer HepB vaccine within 12 hours of birth*	No	1991 <sup>4</sup>	2018 <sup>5</sup>
Administer HepB vaccine and HBIG within 12 hours of birth*	Yes	20185	

https://iournals.sagepub.com/doi/10.1177/00333549231175548 (accessed 12/5/25) https://www.cdc.gov/acip/downloads/slides-2025-09-18-19/02-langer-hep-b-508.pdf (accessed 12/5/25)

52

### **Perinatal Hepatitis B Prevention: History and Progress**

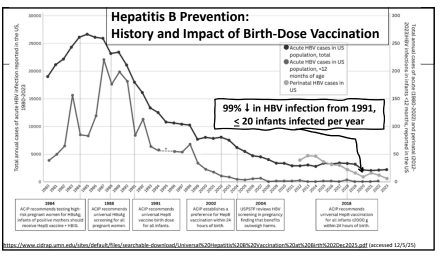
For 2 decades, ACIP has recommended that the first dose of universal infant hepatitis B
vaccination among infants born to HBsAg(-) women occur close to birth:

### ACIP hepatitis B infant vaccination recommendations, United States

Recommendation	Current?	Date of Initial Recommendation	Date Recommendation Superseded or Modified
Infant vaccination strategies			
Universal HepB vaccine before leaving the birth hospital* or within 2 months of age	No	19914	20056
Universal HepB vaccine at the birth hospital*	No	2005 <sup>6</sup>	20185
Universal HepB vaccine within 24 hours of birth*	Yes	20185	

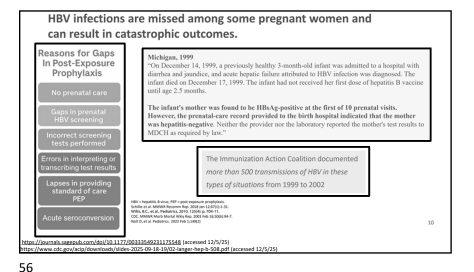
https://journals.sagepub.com/doi/10.1177/00333549231175548 (accessed 12/5/25) https://www.cdc.gov/acip/downloads/slides-2025-09-18-19/02-langer-hep-b-508.pdf (accessed 12/5/25)

53



Why is it important to continue Universal Birth Dose HepB Vaccination?





### Unvaccinated infants remain at risk of non-perinatal HBV acquisition.

- · HBV transmission occurs through percutaneous or mucosal exposure to infectious blood or body
- HBV can remain viable for over 7 days on environmental surfaces at room temperature.1
- Household and Community Transmission: Unvaccinated children living with a person with chronic HBV infection in a household or community setting are at risk for becoming infected.
- Prior to HepB BD, some U.S.-born children born to immigrant mothers without HBV infection had hepatitis B prevalences of 7–11%<sup>2,3</sup> attributable to community or household exposures.
- In the United States, up to 2.4 M people are estimated to have hepatitis B4, and about 50% of people with hepatitis B are unaware of their infection<sup>5</sup>.
- Children who receive HepB BD have higher rates of hepatitis B childhood vaccine series completion and had a positive impact on rates of being up to date for other age-appropriate vaccines 6,7,8

https://journals.sagepub.com/doi/10.1177/00333549231175548 (accessed 12/5/25) https://www.cdc.gov/acip/downloads/slides-2025-09-18-19/02-langer-hep-b-508.pdf (accessed 12/5/25)

57

Rescinding Universal HepB BD vaccination recommendations among infants born to HBsAg (-) women may result in more cases of perinatal HBV infection. **Potential Risks of Rescinding Potential Benefits of Rescinding Universal HepB BD Recommendations Universal HepB BD Recommendations** Increased cases of perinatal HBV transmission Reductions in rare cases of hepatitis B birth dose

vaccination adverse events

Increased administrative complexity and failure points for providers and health systems

Lack of safety net given gaps in access to prenatal care, HBV screening, and HBIG access

Disproportionate harm to patients without insurance or low healthcare engagement

Lower rates of hepatitis B childhood vaccine series

Higher lifetime healthcare costs from missed opportunities to prevent and eliminate hepatitis B

https://journals.sagepub.com/doi/10.1177/00333549231175548 (accessed 12/5/25) https://www.cdc.gov/acip/downloads/slides-2025-09-18-19/02-langer-hep-b-508.pdf (accessed 12/5/25)

## Adverse Effect Assessment for HepB Vaccination...

An independent evidence review of the safety, effectiveness, and public health impact of universal hepatitis B vaccination at birth to compare current recommendations with a delayed first hepatitis vaccine dose at one month or more after birth.





The Vaccine Integrity Project is supported by an unrestricted gift from Alumbra Innovations Foundation to the Center for Infectious Disease Research and Policy (CIDRAP) at the University of Minnesota; no funds from pharmaceutical companies or any other public or private sources were used to support this project. Each member of the research team completed a declaration of interest form, which was reviewed and confirmed by an independent source not related to the project. No members reported a personal of financial interest or relationship related to the content of the report.

https://www.cidrap.umn.edu/sites/default/files/searchable-download/Universal%20Hepatitis%20B%20Vaccination%20at%20Birth%202Dec2025.pdf (accessed 12/5/25)

## **Adverse Effect Assessment for HepB Vaccination...**

### Safety of the hepatitis B birth dose

Results of randomized trials, large national safety monitoring programs, and long-term follow-up studies consistently demonstrate that the hepatitis B vaccine is safe regardless of vaccine timing. No safety benefits were identified for a delayed first dose versus vaccination at birth.

#### Key Summary Findings:

- · Mild-to-moderate short-term reactions:
  - tenderness, redness and swelling at the injection site, fussiness, transient low-grade fever
- No increased incidence of long-term AEs, SAEs, deaths:
  - "...rare deaths following hepatitis B vaccination at birth have been extensively studied and found not to be causally associated with vaccination."
- 4 studies directly compared safety for birth dose and delayed dose:
  - "...no increased risk of any short- or long-term AE or SAE in infants administered the vaccine at birth compared with delayed administration."

https://www.cidrap.umn.edu/sites/default/files/searchable-download/Universal%20Hepatitis%20B%20Vaccination%20at%20Birth%202Dec2025.pdf (accessed 12/5/25)

## **Adverse Effect Assessment for HepB Vaccination...**

### Key Findings:

### Safety of the hepatitis B birth dose

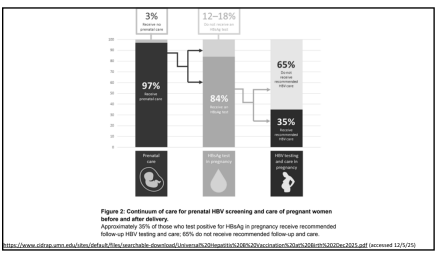
Results of randomized trials, large national safety monitoring programs, and long-term follow-up studies consistently demonstrate that the hepatitis B vaccine is safe regardless of vaccine timing. No safety benefits were identified for a delayed first dose versus vaccination at birth.

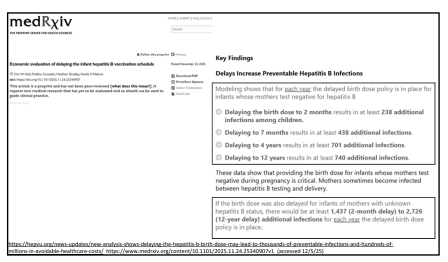
#### Conclusion

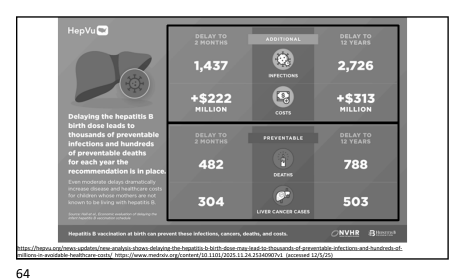
61

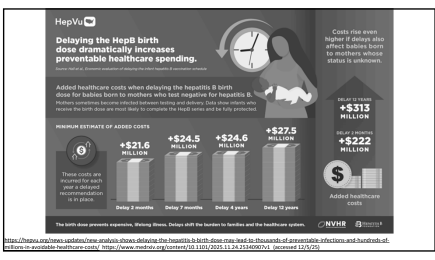
This review found no benefit related to vaccine safety or protection of a delayed first dose compared with vaccination at birth, but identified critical risks of changing current US recommendations.

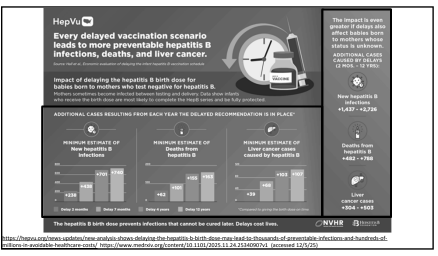
https://www.cidrap.umn.edu/sites/default/files/searchable-download/Universal%20Hepatitis%20B%20Vaccination%20at%20Birth%202Dec2025.pdf (accessed 12/5/25)











## Results of the 12/5/25 ACIP Meeting & Vote Details

#### Vote #1:

- · For infants born to HBsAg-negative women:
  - ACIP recommends individual-based decision-making, in consultation with a health care
    provider, for parents deciding when or if to give the HBV vaccine, including the birth dose. (1)
    For those not receiving the HBV birth dose, it is suggested that the initial dose is administered
    no earlier than 2 months of age. Y/N
    - (1) Parents and health care providers should consider vaccine benefits, vaccine risks, and infection
      risks. Parents and health care providers should also consider whether there are risks, for example,
      such as a household member is HBsAg-positive or when there is frequent contact with persons who
      have emigrated from areas where Hepatitis B is common.

#### Vote Results:

- 8-Yes (Dr. Catherine Stein, Dr. Retsef Levi, Dr. Vicky Pebsworth, Dr. Robert Malone,
   Dr. Hillary Blackburn, Dr. James Pagano, Dr. Evelyn Griffin and Dr. Kirk Milhoan)
- 3-No (Dr. Cody Meissner, Dr. Joseph Hibbeln and Dr. Raymond Pollak)

https://www.cnn.com/health/live-news/cdc-vaccine-meeting-hepatitis-b-12-05-25 (accessed 12/5/25)

## Results of the 12/5/25 ACIP Meeting & Vote Details

#### Vote #2:

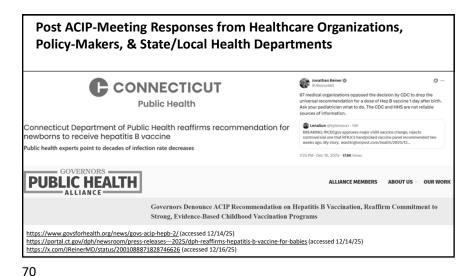
When evaluating the need for a subsequent HBV vaccine dose in children, parents
should consult with health care providers to determine if a post-vaccination anti-HBs
serology testing should be offered. Serology results should determine whether the
established protective anti-HBs titer threshold of ≥10 mIU/mL has been achieved. The
cost of this testing should be covered by insurance. Y/N

#### Vote Results:

- 6-Yes (Dr. Retsef Levi, Dr. Vicky Pebsworth, Dr. Robert Malone, Dr. James Pagano, Dr. Evelyn Griffin and Dr. Kirk Milhoan)
- 4-No (Dr. Cody Meissner, Dr. Joseph Hibbeln, Dr. Raymond Pollak and Dr. Hillary Blackburn)
- 1-Abstained (Dr. Catherine Stein)

https://www.cnn.com/health/live-news/cdc-vaccine-meeting-hepatitis-b-12-05-25 (accessed 12/5/25)





# Selection of Vaccines against Respiratory Viruses (Influenza & COVID-19)



# **Case Example: Seasonal Influenza Vaccine**

- · A 70-year-old male (he/him) comes to the consultation window of your pharmacy
  - He would like to receive a vaccination for the Flu
  - His only chronic health issues are hypertension and hypercholesterolemia (both effectively managed)
- · Which of the following products would be PREFERRED to administer to this patient?
  - a) Inactivated influenza vaccine (IIV3, Afluria)
  - b) High-dose inactivated influenza vaccine (HD-IIV3, Fluzone High-Dose)
  - c) Adjuvanted inactivated influenza vaccine (alIV3, Fluad)
  - d) Live Attenuated Influenza Vaccine (LAIV3, FluMist)

#### **Selection of Influenza Vaccines**

- Main patient considerations for selection of product [not exhaustive list]:
  - Age (above/below 65 y.o.)
  - Immunosuppression / Receipt of Immunosuppressive medications
  - · Pregnancy status
  - Allergic reaction to previous Influenza vaccinations

https://www.cdc.gov/flu/hcp/acip/index.html

## **Selection of Influenza Vaccines**

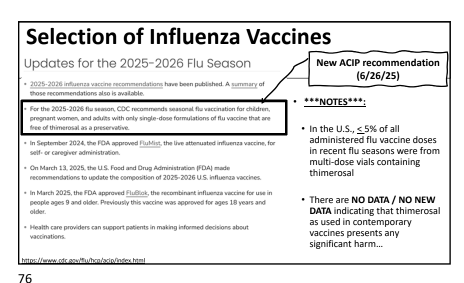
Influenza Vaccine Selection

- Available vaccines, approved ages, and dose volumes are listed in Table 1.
- ACIP recommends all recipients receive seasonal influenza vaccines only in single dose formulations that are free of thimerosal as a preservative (Table 1).
- All persons should receive an age-appropriate vaccine, with the exception that solid organ transplant recipients aged 18 through 64 years who are receiving immunosuppressive medication regimens may receive HD-IIV3 or alIV3 as acceptable options (see Immunocompromised Persons).
- With the exception of Adults Aged ≥65 Years, for whom HD-IIV3, RIV3, and allV3 are preferred (see below), there are no preferences for any specific vaccine when more than one age-appropriate product is available.

https://www.cdc.gov/flu/hcp/acip/index.html

74

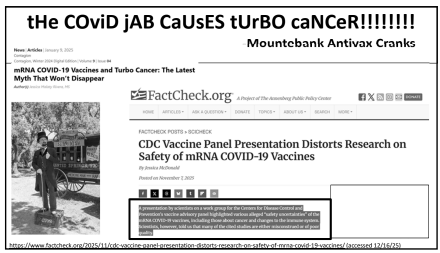
# Adults Aged ≥65 Years ACIP recommends that adults aged ≥65 years preferentially receive any one of the following: High-dose inactivated influenza vaccine (HD-IIV3, Fluzone High-Dose), Recombinant influenza vaccine (RIV3, Flublok), or Adjuvanted inactivated influenza vaccine (aIIV3, Fluad). If none of these three vaccines is available at a vaccination opportunity, then any other age-appropriate influenza vaccine should be used. Data support greater potential benefit of high-dose inactivated, adjuvanted inactivated, or recombinant vaccines relative to standard-dose unadjuvanted IIVs in this age group, with the most data available for HD-IIV3: but comparisons of these vaccines with one another are limited.

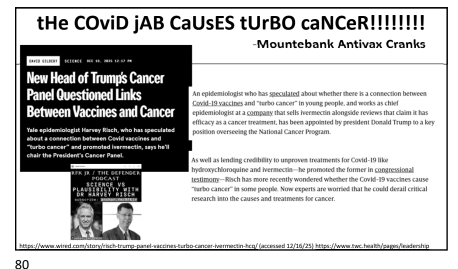


## Case Example: Seasonal Influenza Vaccine

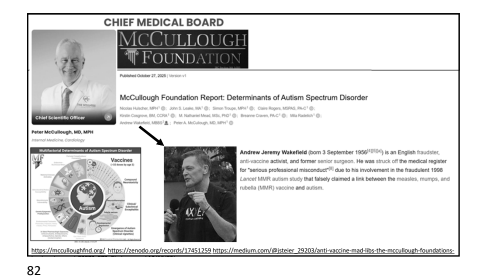
- A 70-year-old male (he/him) comes to the consultation window of your pharmacy
  - He would like to receive a vaccination for the Flu
  - His only chronic health issues are hypertension and hypercholesterolemia (both effectively managed)
- Which of the following products would be <u>PREFERRED</u> to administer to this patient?
  - a) Inactivated influenza vaccine (IIV3, Afluria)
  - b) High-dose inactivated influenza vaccine (HD-IIV3, Fluzone High-Dose)
  - c) Adjuvanted inactivated influenza vaccine (aIIV3, Fluad)
  - d) Live Attenuated Influenza Vaccine (LAIV3, FluMist)

COVID-19 Vaccination









#### A Quick FYI about COVID-19 Vaccination Safety...

COVID-19 mRNA Vaccination and 4-Year All-Cause Mortality Among Adults Aged 18 to 59 Years in France JAMA Netw Open
Published Online: December 4, 2025
2025;8(12):e2546822.
doi:10.1001/jamanetworkopen.2025.46822

• 4-year all-cause mortality in French National Health Data System cohort

• 22.7 million vaccinated, 5.9 million unvaccinated 18-59 year-olds in 2021

Figure. Estimation of All-Cause Mortality at 4 Years in Vaccinated Compared With Unvaccinated Individuals Using Weighted Cox Models: Main and Stratified Analyses

Characteristic	Unvaccinated, No. of events/total No.	Vaccinated, No. of events/total No.	wHR (95% CI)	Lower risk	High risk
Overall	32662/5932443	98429/22767546	0.75 (0.75-0.76)	10	
Age, y					
18-29	2769/1777782	6604/6553907	0.65 (0.62-0.68)		
30-39	4488/1737069	11254/5535008	0.79 (0.76-0.82)		
40-49	8333/1322270	27771/5889066	0.79 (0.76-0.80)	-8	
50-59	17080/1095322	52800/4789565	0.78 (0.77-0.80)		
Sex					
Male	22062/3056404	64946/11078943	0.76 (0.75-0.77)		
Female	10600/2876039	33483/11688603	0.74 (0.73-0.76)		

A Quick FYI about COVID-19 Vaccination Safety...

COVID-19 mRNA Vaccination and 4-Year All-Cause Mortality Among Adults Aged 18 to 59 Years in France

JAMA Netw Open
Published Online: December 4, 2025
2025;8;(12):e2546822.
doi:10.1001/jamanetworkopen.2025.4682

Table 3. RI of Short-Term Mortality, All Causes, by Cancer, External Causes, Circulatory Diseases, and COVID-19, Within 6 Months Following Vaccination, Using Adapted SCCS Models<sup>a</sup>

Table 3. RI of Short-Term Mortality, All Causes, by Cancer, External Causes, Circulatory Diseases, and COVID-19, Within 6 Months Following Vaccination, Using Adapted SCCS Models<sup>a</sup>

	Cause of death, RI	(95% CI)			
Risk window <sup>b</sup>	All-cause	Tumor	Circulatory diseases	External causes	COVID-19
6 mo After dose 1	0.65 (0.63-0.67)	0.71 (0.67-0.76)	0.63 (0.57-0.71)	0.63 (0.58-0.68)	0.73 (0.59-0.91)
6 mo After dose 2	0.76 (0.74-0.79)	0.85 (0.81-0.89)	0.74 (0.66-0.83)	0.78 (0.71-0.86)	0.29 (0.23-0.36)
6 mo After dose 3	0.80 (0.76-0.84)	0.83 (0.77-0.89)	0.76 (0.65-0.88)	0.95 (0.83-1.09)	0.40 (0.30-0.52)
6 mo After any dose	0.71 (0.69-0.73)	0.80 (0.77-0.84)	0.68 (0.62-0.76)	0.67 (0.61-0.72)	0.39 (0.32-0.47)

Abbreviations: RI, relative incidence; SCCS, self-controlled case series.

https://jamanetwork.com/journals/jamanetworkopen/fullarticle/2842305 (accessed 12/15/25)

84

#### A Quick FYI about COVID-19 Vaccination Safety...

Table 2. Comparison of Causes of Death Between Vaccinated and Unvaccinated Individuals up to December 31, 2023, Using Weighted Cox Models Among Those Included in the 4-Year Mortality Study\*

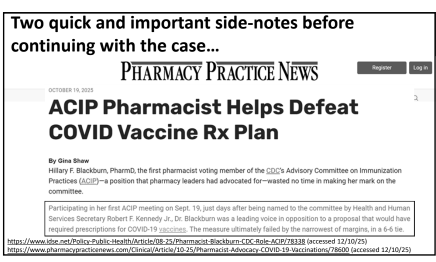
		Incidence per 1 million		Hazard ratio	
ICD-10	Primary causes of death	Among vaccinated	Among unvaccinated	Crude	Weighted
NA	Unknown (unlinkable)	199	327	0.55 (0.52-0.58)	0.58 (0.55-0.61)
A, B	Infectious and parasitic diseases	28	45	0.55 (0.48-0.64)	0.63 (0.54-0.73)
C, D0-D4	Tumors	769	853	0.81 (0.79-0.84)	0.85 (0.83-0.88)
C50 and D05	Including breast cancer	76	103	0.67 (0.61-0.73)	0.68 (0.61-0.74)
C10-C20, D010-D012	Including colorectal cancer	62	66	0.85 (0.76-0.95)	0.89 (0.80-0.99)
C33, C34, D021, D022	Including lung cancer	174	194	0.81 (0.76-0.86)	0.85 (0.79-0.90)
Other codes in C or D0 to D04	Including other cancer	456	491	0.84 (0.80-0.87)	0.89 (0.85-0.92)
D5-D8	Diseases of the blood, hematopoietic organs, and certain immune system disorders	6	13	0.46 (0.35-0.60)	0.50 (0.35-0.68)
U071, U072, U109	COVID-19	18	85	0.20 (0.17-0.23)	0.26 (0.22-0.30)

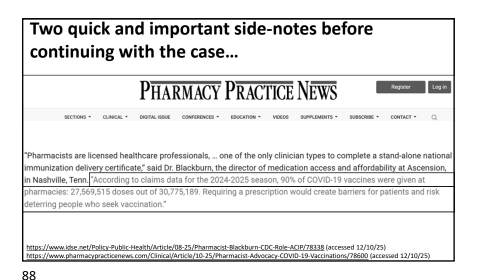
85

https://jamanetwork.com/journals/jamanetworkopen/fullarticle/2842305 (accessed 12/15/25)

## **Case Example: Seasonal COVID-19 Vaccine**

- A 10-year-old girl (she/her) is brought to your pharmacy by her mother; they come to the consultation window of your pharmacy
  - Mom indicates that her daughter is on chronic immunosuppressive medications.
  - She would like her daughter to receive her yearly vaccination for COVID-19
  - You note that she has received a "complete" 3-dose initial COVID-19 vaccine series last year
- · Which of the following vaccine products could be administered to this patient?
  - a) Moderna Spikevax
  - b) Moderna mNexspike
  - c) Pfizer-BioNTech Comirnaty
  - d) Novavax Nuvaxovid
- · What should the administration schedule be?





#### Two quick and important side-notes before continuing with the case...

CDC

89

Morbidity and Mortality Weekly Report (MMWR)

Effectiveness of 2024–2025 COVID-19 Vaccines in Children in the United States – VISION, August 29, 2024–September 2, 2025

Weekly / December 11, 2025 / 74(40);607-614

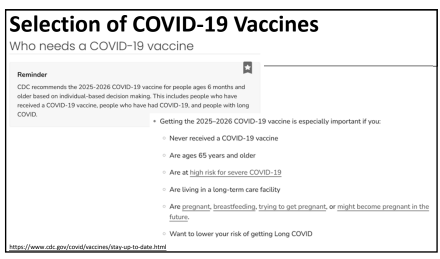
- Vaccine effectiveness (VE) against COVID-19-Associated Emergency Dept/Urgent Care visits:
  - · Children aged 9mo 4 yo: 77% VE (95% CI 62-86%) • Children aged 5mo - 4 yo: 45% VE (95% CI 25-59%)

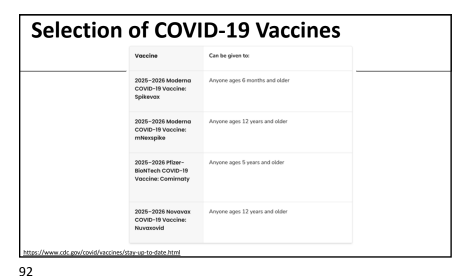
https://www.cdc.gov/mmwr/volumes/74/wr/mm7440a1.htm (accessed 12/14/25) https://www.pharmacypracticenews.com/Clinical/Article/10-25/Pharmacist-Advocacy-COVID-19-Vaccinations/78600 (accessed 12/10/25)

#### **Selection of COVID-19 Vaccines**

- Main patient considerations for selection of product and administration schedule [\*not an exhaustive list]:
  - Age
  - Immunosuppression / Receipt of Immunosuppressive medications
  - Risk of Severe COVID-19 infection
  - Pregnancy

https://www.cdc.gov/covid/hcp/vaccine-considerations/index.html#toc





## Case Example: Seasonal COVID-19 Vaccine

- A 10-year-old girl (she/her) is brought to your pharmacy by her mother; they come to the
  consultation window of your pharmacy
  - Mom indicates that her daughter is on chronic immunosuppressive medications.
  - She would like her to receive her yearly vaccination for COVID-19
  - You note that she has received a "complete" 3-dose initial COVID-19 vaccine series last year
- Which of the following vaccine products could be administered to this patient?

a) Moderna Spikevax

b) Moderna mNexspike

c) Pfizer-BioNTech Comirnaty

d) Novavax Nuvaxovid

What should the administration schedule be?

## Case Example: Seasonal COVID-19 Vaccine

- A 10-year-old girl (she/her) is brought to your pharmacy by her mother; they come to the consultation window of your pharmacy
  - Mom indicates that her daughter is on chronic immunosuppressive medications.
  - She would like her to receive her yearly vaccination for COVID-19
  - You note that she has received a "complete" 3-dose initial COVID-19 vaccine series last year
- Which of the following vaccine products could be administered to this patient?
  - a) Moderna Spikevax
  - b) Moderna mNexspike
  - c) Pfizer-BioNTech Comirnaty
  - d) Novavax Nuvaxovid
- What should the administration schedule be?

#### **Selection of COVID-19 Vaccines**

COVID-19 Vaccination Guidance for People Who Are Immunocompromised

AT A GLANCE

- COVID-19 vaccination is recommended for people ages 6 months and older who are moderately or severely immunocompromised based on individual-based decision-making (also known as shared clinical decision making).
- There is a modified COVID-19 vaccination schedule for people who are moderately or severely immunocompromised.
- \* People can self-attest to being moderately or severely immunocompromised and receive COVID-19 vaccination.
- Administering COVID-19 vaccines should not be delayed in patients taking immunosuppressive therapies.

https://www.cdc.gov/covid/hcp/vaccine-considerations/immunocompromised.html

## Selection of COVID-19 Vaccines Table 2: 2025–2026 COVID-19 vaccination schedule for people who are moderately or severely immunocompromised, November 4, 2025 2b: Ages 5–11 years NOTE Moderna (Spikevax) and Pfizer-BioNTech vaccines are approved for this age group. In Table 2b, Moderna refers to Spikevax. See footnote\* for guidance on children who transition from age 4 years to age 5 years during the initial vaccination series.

96

#### **Selection of COVID-19 Vaccines** COVID-19 vaccination history before Number of 2025–2026 Recommended 2025–2026 vaccine<sup>‡</sup> and interval between doses 2025-2026 vaccine doses indicated Completed the 3-dose initial series before 2025–2026 vaccine: Administer 2 doses of 2025–2026 vaccine spaced 6 months apart 2025–2026 Dose 1 (Moderna or Pfizer-BioNTech): At least 8 3 or more doses Moderna or 3 or more doses Pfizer-BioNTech 2025-2026 Dose 2 (Moderna or Pfizer-BioNTech): 6 months minimum interval 2 months) after 2025–2026 Dose 1 Children who transition from age 4 years to age 5 years during the initial vaccination series should complete the 3-dose series using the dosage for children ages 5–11 years for all doses received on or after turning age 5 years: - Moderna series: 2025–2026 Moderna, 0.25 mL/25 ug; there is no dosage change - Pfizer-BioNTech series: 2025-2026 Pfizer-BioNTech, 0.3 mL/10 ug COVID-19 vaccination history refers to all doses of COVID-19 vaccine from any manufacturer received before the availability of the 2025–2026 COVID-19 Dosage for Moderna (Spikevax): 0.25 mL/25 ug; dosage for Pfizer-BioNTech: 0.3 mL/10 ug. https://www.cdc.gov/covid/hcp/vaccine-considerations/immunocompromised.html

## Case Example: Seasonal COVID-19 Vaccine

- A 10-year-old girl (she/her) is brought to your pharmacy by her mother; they come to the
  consultation window of your pharmacy
- Mom indicates that her daughter is on chronic immunosuppressive medications. She would like her to receive her yearly vaccination for COVID-19
- You note that she has received a "complete" initial COVID-19 vaccine series in the past year
- Which of the following vaccine products could be administered to this patient?

a) Moderna Spikevax
b) Moderna mNexspike
c) Pfizer-BioNTech Comirnaty
d) Novavax Nuvaxovid

98

What should the administration schedule be?

registrate for 2 route details brief horize up 222 - 2225 Modelle and 2 route details brief horize up 2225 - 2225 Modelle and 2 route details and 2 route details and 2 route details and 2 route details and 2 route at 10 ro

#### **Post-Test Time!**

- The U.S. Dept. of HHS, the CDC, the FDA, and the Advisory Committee on Immunization Practices (ACIP) have been infiltrated by Mountebanks, Antivax Cranks, Grifters, Charlatans, & Toadies...
- Which **ONE** of the following is **NOT** a recommendation that has been **APPROVED** by ACIP in 2025?
  - All adults receive seasonal influenza vaccines only in single dose formulations that are free of thimerosal as a preservative
  - State and local jurisdictions should require a prescription for the administration of a COVID-19 vaccination
  - COVID-19 Vaccination should be based on individual-based decision-making (Shared Clinical Decision Making) for Adults 65 and older
  - d) Universal Hepatitis B Vaccination at birth is no longer recommended for all newborns
  - e) For Hepatitis B vaccination, post-vaccine serology results should determine need for subsequent doses (in the 3-dose series)

## Conclusions, Final Thoughts, and Personal Expert Opinions...

- Based on what appear to be "vibes", the current group of Mountebanks at HHS, FDA, CDC, ACIP have initiated a dangerous erosion of logical U.S. public health policies that had been based on decades of high-quality scientific evidence
- In the very near future, it is very likely that we will have to start suboptimally treating various vaccine-preventable infectious diseases that were once considered "eradicated" or "rare" in the U.S.
- We as Pharmacists can play an important and significant role in:
  - The "Shared Decision Making" for Influenza and COVID-19 vaccination for patients
  - Countering medical and pharmaceutical misinformation currently emanating from oncerespected and trusted U.S. Government Health Agencies

99

## Conclusions, Final Thoughts, and Personal Expert Opinions...

Vaccine RCT spreadsheet aims to show the data, dispel myths about vaccines

Chris Dall, MA, June 18, 2025

- "Living Google Document" developed/maintained by Dr. Jake Scott (Infectious Diseases MD @ Stanford U.)
  - List and summary of ALL RCTs ever conducted for licensed vaccines
  - <u>Link:</u>
     https://docs.google.com/spreadsheets/u/0/d/1bX4SAJwMUufNAkBplhKHOle4
     gdl0BeRhpAXM5hpfV Y/htmlview?pli=1#gid=0

https://www.cidrap.umn.edu/adult-non-flu-vaccines/vaccine-rct-spreadsheet-aims-show-data-dispel-myths-about-vaccines (Accessed 12/14/25)

### Thank you!!!

• Questions???

102

**SESSION CODE:**