Occupational Health Update: Acute Care Facilities 4/23/24

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Disclosures

- No financial relationships to disclose
- No off-label or investigational use of medications and/or devices
- The information and views set out in this presentation are those of the author and do not necessarily reflect the official opinion of the University of North Carolina at Chapel Hill or UNC Health





Objectives ACIP Updates Vaccines for HCPs (Pre-exposure prophylaxis) Post-exposure prophylaxis (Bloodborne Pathogens) COVID-19 Employee Well-Being Civic Health

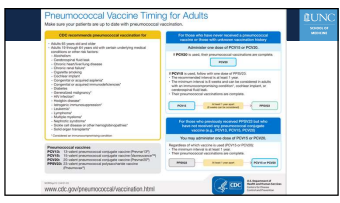




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ACIP January 2022 Update Preumococcal Vaccines New availability of PCV 15 (Merck Sharp & Dohme Corp.) and PCV 20 (Wyeth Pharmaceuticals LLC) Recommendation for PCV20 alone or PCV15 in combination with PPSV23 for previously unvaccinated 65y+ or 19-64 y/o with increased risk If PCV15 is used, minimum interval of 8 weeks until PPSV23 administration Prior PPSV23 – either PCV15 or PCV20 can be used Prior PCV13 – continue with current schedule for PPSV23 afterwards (still under evaluation of added benefit of PCV15 or PCV20 after)



ACIP April 2022 Update

- Hepatitis B Vaccines are now universally recommended for all adults aged 19 – 59 years old instead of based solely on risk factors. This reflects the rising cases of Hepatitis B since nadir in 2014, and acknowledges that risk-based intervention misses people reluctant to disclose.
- Also note that ACIP recommendations for Hepatitis B screening was updated in March 2023 to include testing at least once per lifetime in addition to risk factor based testing

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JYNNEOS for Monkeypox Two vaccines (JYNNEOS and ACAM2000) for orthopoxviruses (including MPX and smallpox). JYNNEOS w/ much less contraindications. Pre- or post- exposure prophylaxis indications based on risk factors (generally intimate, prolonged contact) Most healthcare workers do not need to get this vaccine. Exceptions include HCPs w high risk exposure (caring for +nt for prolonged period)

- include HCPs w high risk exposure (caring for +pt for prolonged period without PPE) and lab personnel handling specimens (current wording as of 4/10/23: "You work in settings where you may be exposed to mpox:
 - You work with orthopoxviruses in a laboratory

ACIP June 2022 Update

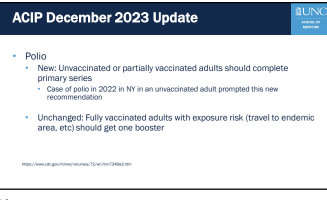
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- You are part of an orthopoxvirus and health care worker response team"
- /www.cdc.gov/mmwr/volumes/71/wr/mm7122e1.htm Vaccines |https://www.cdc.gov/poxvirus/mpox/vaccines/index.html Mpox | Poxvirus | CDC

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ACIP June 2023 Update RSV Vaccine (Abrysvo or Arexvy) for adults 60+ (shared decision making) Single dose (for now), high efficacy over two RSV seasons Can be coadministered with other vaccines Abrysvo is also recommended for pregnant people 32 - 36 wks GA from Sept - Jan When vaccinating adults 60+ years, it should be done year round (in contrast with pregnant people and babies only during RSV season)

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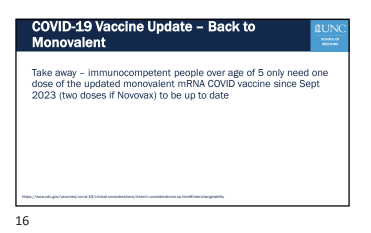




Personnel

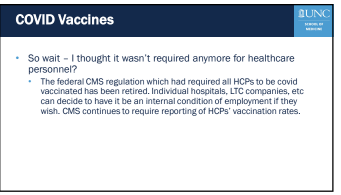
HCP Vacci	nation Recommendations					
Vaccination	Recommendation					
COVID-19	If not up to date, provide COVID-19 vaccine. Pts >6 months should receive 1 bivalent booster. (For adults, easiest to ask if they've received a booster since Sept 1, 2022.)					
Hepatitis B	If no prior dose, either 2 doses of Heplisav-B or 3-dose series of either Engerix or Recombivax Obtain serology 1-2 months after final dose					
Influenza	Give 1 dose annually					
MMR	HCP born in 1957 or later need 2-doses of MMR, 4 weeks apart if no prior immunity or vaccination. Before 1957, consider serology testing and dosing if needed					
Varicella	If no prior infection, serologic immunity, prior vaccination, give 2 doses of varicella vaccine 4 weeks apart					
Tetanus, diphtheria, pertussis	Give 1 dose to all who have not received previously. Each pregnancy. Booster every 10 years (Td or Tdap)					
Meningococcal	Routinely to microbiologists exposed to isolates of N. Meningitidis					
https://www.cdc.gov/va	ccines/adults/rec-vac/hcw.html					

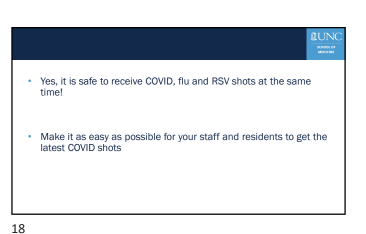
	nper	ent)					
COVID-19 vaccination history prior to updated (20032024 Formula) vaccine*	Updated (2023-2024 Formula) vaccine	Number of updated (2023-2024 Formula) down indicated	Donage prot Aug	Vaccine vial Cap and label colors ⁶	interval between dataets		
Uniacciruited	Moderna	1	0.5 mL/50 ug	Dark blue cap; blue libel			
1 or more doses any millity, 1 or more doses any millity, 1 or	ot						
	Noveman	2	0.5 mL/5 ug r5 protein and 50 ug Matrix M adjuvant	Blue cap; blue label	Dose 1 and Dose 2:3-8 weeks		
	OR						
	Pfizer BioNillech	а.	0.3 mL/30 vg	Gray cap: 37 av Label			
	Moderna	3	0.5 mL/50 kg	Dark blue cap: blue label	At least 8 weeks after last dose		
including in combination with any Original monovalent or bivalent	OR						
COMD-19 vaccine doses	Tepravas	3.	0.5 mL/5 ug r5 protein and 50 ug Matrix M adjuvant	Dive cap: blue label	At least 8 weeks after last close		
	OR						
	Pfeer-BioNTech	2	0.3 mi/30 ug	Gray cap: gray Label	At least 8 weeks after last dose		

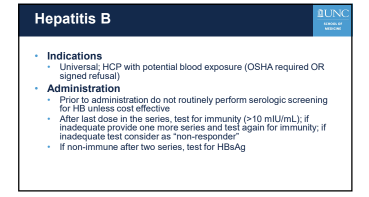


ACIP COVID-19

Vaccine





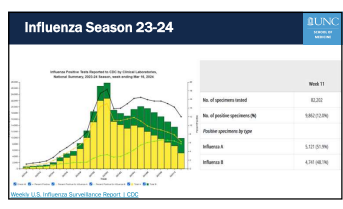


Hepatitis B HEPLISAV-B approved in late 2017 • Adults > 18 years of age • Two doses one month apart Not studied in hemodialysis patients Table 7 Study 3: Seroprotection Rates of HEPLISAV-B and Engerix-B* (ages 18 - 70 years) Age (years) HEPLISAV-B^a Engerix-B^a (HEPLISAV-B minus Engerix-B Difference (95% CI) 6.1% (2.8, 12.6)* N SPR (95% CI) 174 100.0% (97.9, 100.0) SPR (95% CI 93.9% (87.3, 97 18-29 30-39 40-49 99 326 92.0% (88.5, 94.7) 518 84.2% (80.7, 87.2) 79.7% (76.6, 82.5) 632 98.9% (97.7, 99.6) 974 97.2% (96.0, 98.2) 6.9% (4.2, 10.4) 13.1% (9.9, 16.6)*
 50-59
 1439
 95.2% (94.0, 96.3)
 758
 79.7% (76.6, 82.5)

 60-70
 1157
 91.6% (89.9, 93.1)
 588
 72.6% (68.8, 76.2)
 15.5% (12.6, 18.7)*

19.0% (15.2, 23.0)

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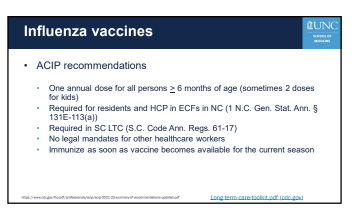




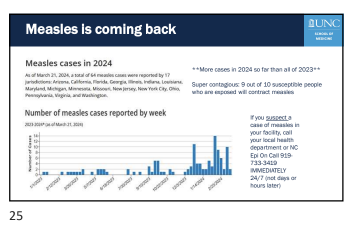
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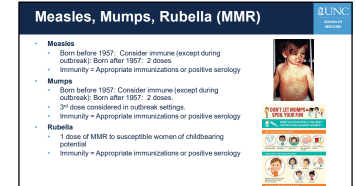


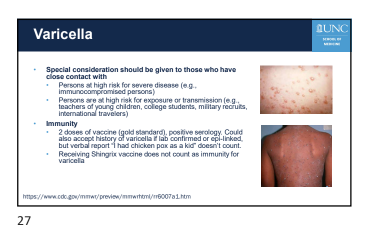


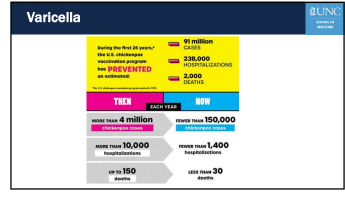


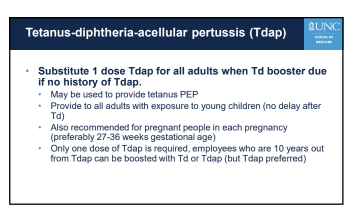


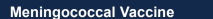












- Recommended for adults had high risk of disease (persistent complement deficiency, functional or anatomic asplenia, or HIV infection (adolescents)).
 Two vaccines series are needed: MenACWY and Serogroup B (MenB)
- MenACWY
- Immunosuppressed 2 doses of MenACWY and boosters every 5 years, 2 or 3-dose MenB
 Microbiologists – 1 dose, booster every 5 years (MenACWY), 2 or 3dose MenB
- Now they could get the combo MenABCWY vaccine when both are indicated Anatomic/functional asplenia patients should be vaccinated against MenACWY/MenB

Tuberculosis Surveillance







Tuberculin Skin Test Conversions and Oc	upational
Exposure Risk in US Healthcare Workers	
Claudia C. Dobler, ¹³ Wingdon H. Farah, ² Monar Alsonvas, ¹ Khaled Mohammed, ²³ Lawa E. Breeher, ¹ M. Hassan ''Dision of Poestris, Orząpzional and Aeropans Medicine and 'Tothera-Band Pacino Gente, Mayo Dani, Rubent, Mareant, Momenta, Mareapin	
Bedgraund, Henhner wecker (HCW) in aderge occupational thereations screening of contracting thereadows in the workpoice in a string with a low bodynomi baberuid exilate the risk of tabercalini skin tori (TST) conversion and the risk of ecopyrisms at herea- tion and the string of the string screening of the string of ecopyrisms (a large transport of the string of the string screening screening screening screening screening screening and the string screening during the storing screening screening screening screening screening screening during the string screening baryons, and the screening screening screening screening screening screening baryons at the screening scr	s incidence is unclear. We aimed to toois infection among HCWs in such a centric in the US Midwest who had Only 9 (7%) of the converters had a majority of TST converters (66%) had mess setting, we demonstrated an inguiret indexeduct cases. In this

Baseline (preplacement) screening and testing. All U.S. health care personnel should have baseline TB screening, including an individual risk assessment, which is necessary for interpreting any test result. IGRAs (quant gold or T spot) or tb skin tests can be used. Follow CDC algorithm for interpretation. Serial screening and testing for health care personnel without LTBI is NOT indicated. In the absence of known exposure or evidence of ongoing TB transmission, U.S. health care personnel (as identified in the 2005 guidelines) without LTBI should not undergo routine serial TB screening or testing at any interval after baseline (e.g., annually.) Could consider annual screening with high risk groups like respiratory therapists. Health care personnel with LTBI and no prior treatment should be offered, and strongly encourage to complete treatment with a recommended regimen, including short-course treatments, unless a contraindication exists

Bloodborne Pathogens



Bloodborne Pathogens

- Approximately 385,000 needle sticks and other sharps-related injuries to hospital-based healthcare personnel each year.
- 58 total known occupationally acquired HIV cases in HCPs; all but 1 were prior to 1999.
- 88% (50/57) of the documented cases of occupational HIV transmission from 1985-2004 involved a percutaneous exposure. Of those, 45/57 involved a hollow-borne needle.
- 41% of sharp injuries occur during use; 40% after use/<u>before</u> disposal; 15% during/after disposal

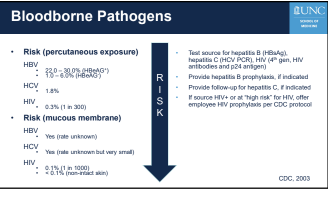
https://www.cdc.gov/mmwr/preview/mmwrhtml/mm6353a 4.htm

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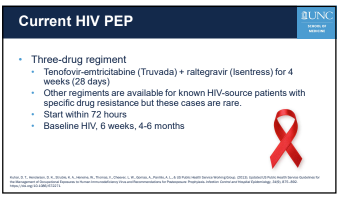
OSHA Bloodborne Pathogens Standard Employers must establish a written exposure control plan and ٠ provide annual training Mandates use of universal precautions (all body fluids assumed contaminated except sweat) Employers must utilize engineering and work practice controls • to minimize/eliminate exposure (e-CFR 1910.1013) https://www.osha.gov/laws-regs/regulations/standardnumber/1910/1910.1030 40

NUN OSHA Bloodborne Pathogens Standard Requires offering hepatitis B vaccine to persons with the potential for exposure Testing of exposed employees for Hepatitis B and HIV Post-exposure prophylaxis must be immediately available as per CDC guidelines All work-related needle stick injuries and cuts from sharp objects that are contaminated with another person's blood or other potentially infectious material are OSHA-reportable <u>regardless of</u> the source patient disease status. (e-CFR 1910.1013)



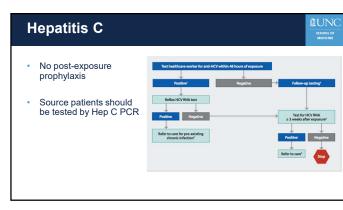
kposi							
Infection Status of Source Patient	Baseline Labs	2 Weeks	4 Weeks	6 Weeks	4 Mantha	6 Months	
DATE: >							
HIV positive	HIV test - 4 th generation	Lab - only if baseline abnormal or dinical indication		HQV test - 4" generation	HIV test - 4" generation		
Hillsåg positive	If source positive and HCP unknown, need HBaAb. If HBaAb ≥12 mDU/mL. testing complete. If HBAb <12 mDU/mL, need anti- HBC & HBaAg at baseline					• Anti-HBc • HBsAg	
Hepatitis C RNA PCR positive	Anti-HCV (Hepatitis C antibody)	Lab - only if baseline abnormal or dinical indication		HCV RNA PCR	Anti-HCV (Hepatitis C antibody)		
Unknown source	HEV test – 4° generation Brounce unknown and HEP Heakbu witknown, need Heakb. If Heakb J 22 mit/Unit, testing complete. If Heakb <12 mit/Unit, need artD- HEC & https/g at baseline HEV artspory	Lab - only if baseline abnormal or dinical indication		Infly test - 4 th generation HCV RNA PCR	HIV test - 4 ^e generation Anti-HCV (Hepatitis C antibody)	+ Anti-Hite + HiteAg	

		MEDICINE
occupational health provider shall notify the healthcare pr exposure has occurred. This healthcare provider sha	wown, the attending physician or responsible for the exposed perso ovider of the source case that an all arrange HIV testing of the source HIV+) and notify the OHS provider	e



 Universal; HCP with potential blood exposure (OSHA require or HCP may decline) No need to routinely obtain Hep B titers if an employee has documented vaccine series and a positive titer In practice, we usually titer and give a booster if titer is < 10 mIU/m For known non-responders, with exposure they should get Hepatiti B Immune Globulin (HBIG) within 24 hours (up to 7 days after exposure) 	L

Нер	atiti	s B		Postexposure Management of Health Care Personnel after Occupational Exposure to Blood and Body Fluids, by Health Care Personnel HepB Vaccination and Response Status						
				HepB Vaccination and Response Status	Postexposure testing results for source patient (HBsAg)	Postexposure testing results for HCP (anti-HBs)	HBIG* postexposure prophylaxis	Vaccination postexposure prophylaxis	Postvaccination Serologic Testing'	
				Documented responder ⁴ after complete series (3 or more doses)	No action needed	No action needed	No action needed	No action needed	No action needed	
	HBsAg	Anti-HBc	HBsAb*	Documented nonresponder* after 2 complete	Positive/ unknown		2 doses HBIG separated by 1 month	No action needed	No action needed	
Acute infection	Positive	IoM positive	Negative	series	Negative	No action needed	No action needed	No action needed	No action needed	
Infection resolved	Negative	IgG Positive	Positive		Positive/unknown	less than 10 mlU/mL**	1 dose HBIG	Initiate	Yes	
Chronic infection	Positive	IgG Positive	Negative	Response unknown after a complete series	Negative	less than	None	Initiate	Nex.	
Vaccinated	Negative	Negative	Positive		-	10 mIU/mL		revectination		
Susceptible	Negative	Negative	Negative		Any result	greater than or equal to 10 mIU/mL	No action needed	No action needed	No action needed	
Olera, Willam, Parga, Julán, & Gastelsbonda, Johanna. (2018). Serology of hepatitis B virus: multiple aconarios and multiple econes. <i>Revisita colonibiana</i> de Gastroenterologie, 33(4), 411–622. <u>https://doi.org/10.22510/25007440.327</u>			Unwaccinated/ incompletely vaccinated or vaccine refusers	Positive/ unknown	**	1 dose HBKS	Complete vaccination	Yes		
				Negative	No action needed	None	Complete vaccination	Yes		
				"HBIG should be adm administered greater should be administered	than 7 days after perce	rly as soon as possible staneous, mucosal, or i ic injection sites.	after exposure when nonintact skin exposu	indicated. The effectiv res is unknown. HBIG	eness of HBIG when and Hep8 vaccine	
				Should be performed to avoid detection of tration of anti-HBs (gr	passively administered	ne last dose of the Hep Lanti-HBs) using a qua 10 miU/ml.).	8 vaccine series (and ntitative method that	4 to 6 months after ac allows detection of th	Iministration of HBIG reprotective concen-	
						i-HBs greater than or e				
				¹ A nonresponder is de						
				source patient who is	H8sAg-positive or has re and follow-up testin	a approximately 6 mo	s, should undergo ba	seline testing for HRV	infection as soon as	



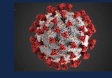
Follow-up Testing

- Hepatitis B
- Not required if employee has immunity
- HIV

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- Dependent on source patient and available testing
- Hepatitis C
 - · Dependent on source patient, test for HCV antibodies and HCV RNA

Occupational Health and COVID



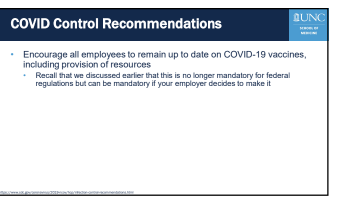


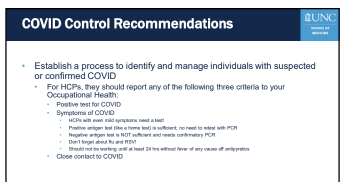
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COVID Control Recommendations Updated May 8, 2023 Encourage all employees to remain up to date on COVID-19 vaccines, including . provision of resources Establish a process to identify and manage individuals with suspected or confirmed $\ensuremath{\mathsf{COVID}}$ Implement source control measures (changed from earlier recommendations) • Implement universal use of personal protective equipment for HCP Optimize use of engineering controls and indoor air quality . Perform SARS-CoV-2 viral testing • Create a process to respond to COVID exposures among HCP and others . 52





COVID-19+ HCP Return to Work

HCP with mild to moderate illness who are not moderately to severely immunocompromised could return to work after

the following criteria have be

- · At least 7 days have passed since symptoms first appeared if a negative viral test* is obtained within 48 hours prior to returning to work (or 10 days if testing is not performed or if a positive test at day 5-7), and
- At least 24 hours have passed since last fever without the use of fever-reducing medications, and
- · Symptoms (e.g., cough, shortness of breath) have improved.

*Either a NAAT (molecular) or antigen test may be used. If using an antigen test, HCP should have a negative test obtained on day 5 and again 48 hours later

HCP who were asymptomatic throughout their infection and are not moderately to severely immunocompromised could return to work after the following criteria have been met:

- At least 7 days have passed since the date of their first positive viral test if a negative viral test* is obtained within 48 hours prior to returning to work (or 10 days if testing is not performed or if a positive test at day 5-
- *Either a NAAT (molecular) or antigen test may be used. If using an antigen test, HCP should have a negative test obtained

on day 5 and again 48 hours later

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What about guarantines for exposures? Work restriction is not necessary for most asymptomatic HCP following a higher-risk exposure, regardless of vaccination status. Examples of when work restriction may be considered include · HCP is unable to be tested or wear source control as recommended for the 10 days following their exposure; HCP is moderately to severely immunocompromised;

- HCP cares for or works on a unit with patients who are moderately to severely immunocompromised, HCP works on a unit experiencing ongoing SARS-CoV-2 transmission that is not controlled with initial interventions;

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Asymptomatic HCPs w COVID exposures Following a higher-risk exposure HCP should:

Have a series of three viral tests for SARS-CoV-2 infection.

- Testing is recommended immediately (but not earlier than 24 hours after the exposure) and, if negative, again 48 hours after the first negative test and, if negative, again 48 hours after the second negative test. This will typically be at day 1 (where day of exposure is day 0), day 3, and day 5.
- Due to challenges in interpreting the result, testing is generally not recommended for asymptomatic people who
 have recovered from SARS-CoV-2 infection in the prior 30 days. Testing should be considered for those who
 have recovered in the prior 31-90 days; however, an antigen test instead of NAAT is recommended. This is because some people may remain NAAT positive but not be infectious during this period.
- · Follow all recommended infection prevention and control practices, including wearing well-fitting source control, monitoring themselves for fever or symptoms consistent with COVID-19, and not reporting to work when ill or if testing positive for SARS-COV-2 infection.
- Any HCP who develop fever or <u>symptoms consistent with COVID-19</u> should immediately self-isolate and contact their established point of contact (e.g., occupational health program) to arrange for medical evaluation and testing.

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Employee Well-being Could be its own lecture Taking good care of employees benefits all: patients, employees, and the business (safer environment, lower turnover, less staffing shortages) Physical and mental well-being Living wages and robust benefits Parental leave Comprehensive DEI (diversity, equity and inclusion) trainings and meaningful reflections in workplace policies/practices, not just lip service Safety from workplace violence Fair PTO policies that disincentivize presenteeism Access to resources for burnout, moral injury







