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Vaccine Adherence: Addressing Myths and Hesitancy

Determining which vaccines are appropriate for your patient is based on several factors (e.g., age, health conditions, lifestyle). Patient fears, myths, and scheduling may be barriers to vaccine adherence. Use this checklist to improve vaccination rates, increase adherence, and overcome barriers.

Goal	Suggested Approach
Identify candidates	Ask about vaccine history. For example, you can ask: "Which vaccines have you received?" "When was your last tetanus shot?" Use these tools to stay current on available vaccines and the latest recommendations for all age groups: Use these tools to stay current on available vaccines and the latest recommendations for all age groups: Use these tools to stay current on available vaccines and the latest recommendations for all age groups: Use these tools to stay current on available vaccines and the latest recommendations for all age groups: Use these tools to stay current on available vaccines and the latest recommendations for all age groups: Use these tools to stay current on available vaccines/ehdules/index.html. There are also online quizzes to determine needed vaccines: Adults: What Vaccines Do You Need? (https://www2.acdc.gov/njc/adultimmsched/). Children and adolescents: What Vaccines Does Your Child Need? (https://www2.cdc.gov/vaccines/childquiz/) Canada: https://www.canada.ca/en/public-health/services/provincial-territorial-immunization-information.html When available, review immunization registry data to determine which vaccine(s) a patient may need. Develop strategies to identify eligible patients. Consider patient ages and chronic medical conditions. For example: Help parents stay on track with childhood vaccinations for infants and young children. Adolescents may need the human papilloma virus (HPV) and meningitis vaccines. Elderly patients may be candidates for the pneumococcal or zoster vaccines. Elderly patients may be candidates for the pneumococcal or zoster vaccines. Adolescents may need a pneumococcal vaccine. Make sure ALL patients six months and older, including pregnant women, receive a flu vaccine yearly. Be familiar with and follow policies for giving vaccines to minors with and without parental consent. US: individual state laws can be found at https://www.vaxteen.org/consent-laws-by-state. Canada: check for provincial age of consent requirements, as ages may vary am
Address hesitancy	 □ Ask about vaccine hesitancy. For example, you can ask, "What keeps you or your child from getting a recommended vaccine?" □ Infants: Ease fears about the number of vaccines infants receive at one time. Evidence suggests that a healthy child's immune system will NOT be damaged or overwhelmed by receiving multiple vaccines at once.³ □ Adolescents: Reassure that the HPV vaccine does NOT increase sexual promiscuity or sexually related outcomes (e.g., pregnancy).⁴ □ Adults: Educate that vaccines not only prevent infections, but also significant infection-related complications. ○ For example, the flu vaccine lowers the risk of flu-related complications (e.g., hospitalizations).⁵

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Ease fears about unfounded myths	Ask about fears and questions. For example, you can ask, "What fears or questions do you have because of things you have heard about vaccines?"
difficultation in the second s	Remind patients that the flu vaccine may cause mild malaise or flu-like symptoms, but it does NOT cause the flu. Tell patients that they can't believe everything they see on the internet about vaccines, as some of the information is false. But reassure them that studies consistently show that vaccines (even old ones that had thimerosal) DO NOT cause autism. Some prefer natural immunity over vaccines. It is not worth the risk, especially for some infections. Stress the risks and complications of disease. For example Severe allergic reactions to the measles, mumps, and rubella (MMR) vaccine occur in about 1 in 1,000,000 doses. But, about one in 1,000 patients infected with measles will die. In adults, data suggest that COVID-19 vaccine-induced immunity protects against reinfection five times better than a previous COVID-19 infection.
Improve adherence	Use strong endorsements. Consider using an "opt-out" approach instead of an "opt-in" approach. Some data suggest proactively scheduling appointments for patients (opt-out approach) to receive a vaccine increases vaccination rates compared to notifying patients that vaccination appointments can be made (opt-in approach). Personalize the conversation. Share that you vaccinate your kids. Ask if they were vaccinated when they were young. In the US, encourage booking future vaccine doses with the first dose. Enroll patients in reminder programs (e.g., calls, texts). In Canada, follow school vaccination programs (where available) to ensure required vaccines are received on schedule. Suggest coordinating care with other providers who offer vaccines (e.g., pharmacies, other medical appointments).

Users of this resource are cautioned to use their own professional judgment and consult any other necessary or appropriate sources prior to making clinical judgments based on the content of this document. Our editors have researched the information with input from experts, government agencies, and national organizations. Information and internet links in this article were current as of the date of publication.

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Vaccinating Immunocompromised Patients

full update September 2024

Concerns are raised when a potentially immunocompromised (i.e., immunosuppressed) patient presents for vaccination. The concern with live vaccines is that the patient might contract the disease from the vaccine. Inactivated vaccines cannot cause disease, and some inactivated vaccines are especially recommended for immunocompromised patients. However, depending on the patient's degree of immunocompromise, response to some vaccines may be suboptimal. For some disease states/vaccinations, titers could be used to assess response. It is important to assess the patient's degree of immunocompromise when making vaccine decisions, especially for live vaccines. When in doubt, consult the specialist caring for the patient's immunocompromising condition.³ If possible, ensure that patients are vaccinated with routine adult vaccinations (plus any others that are specific to their condition) before immunocompromise. And keep in mind that several live vaccines have inactivated alternatives (influenza, typhoid, polio).

--Information in chart may differ from product labeling.--

For help identifying which vaccines are LIVE and which are INACTIVATED, see:

- Vaccines Licensed for Use in the United States at https://www.fda.gov/vaccines-blood-biologics/vaccines/vaccines-licensed-use-united-states.
- Contents of Immunizing Agents Authorized for Use in Canada at https://www.canada.ca/en/public-health/services/publications/healthy-living/canadian-immunization-guide-part-1-key-immunization-information/page-15-contents-immunizing-agents-available-use-canada.html#p1c14t1.

Clinical Question	Pertinent Information of Resource
WHO is or might be	• Patients with cancer affecting the bone marrow or lymphatics. ³
immunocompromised	• Patients being treated with chemo (e.g., alkylating agents, antimetabolites) or radiation , and for three months
in the context of	afterward. 1,3
vaccination?	• Patients receiving immunosuppressive biologics (e.g., anti-TNF agents, lymphocyte-depleting agents). 1,3
	• Patients with complement deficiency, or receiving complement inhibitors (e.g., eculizumab). ^{2.3}
	• Transplant patients. ^{2,3}
	• Patients with congenital (primary) immunodeficiency. 1,3
	• Patients receiving large doses of corticosteroids (see footnote a).
	• HIV patients. Degree of immunocompromise varies widely; consider CD4 count and CD4 precentage.
	• Patients taking immunosuppressants (e.g., high-dose methotrexate, azathioprine, or 6-mercaptopurine doses [see
	footnote a]; calcineurin inhibitors). ³
	Asplenia (increased risk of fulminant bacteremia). ^{2,4}
	• Chronic renal disease. ²

Clinical Question	Pertinent Information of Resource
Can patients with	Also see separate section on immunosuppressive MEDICATIONS , below.—
immunocompromise	Non-live vaccines include killed whole-organism, recombinant, subunit, split-virus, toxoid, polysaccharide,
receive non-live	polysaccharide protein-conjugate, and mRNA vaccines. ^{2,15}
vaccines?	 Because non-live vaccines cannot replicate, they are safe for immunocompromised patients. ¹⁻³ However, these patients may not respond as well as immunocompetent patients. ^{1,3} Consider the following: If risk of infectious exposure is low, consider delaying inactivated vaccines until the person is less immunosuppressed.³ Review vaccination history and administer any needed vaccines at least two weeks before planned immunosuppression to optimize response.³ All vaccines are likely effective in patients with chronic kidney disease, primary complement deficiency, certain phagocytic deficiencies, and nonsevere antibody deficiency (e.g., IgA, IgG subclass).² For information on efficacy in other disease states, see reference 2. Some inactivated vaccines are especially encouraged in immunocompromised patients. For recommendations for specific disease states or conditions (e.g., HCT, solid organ transplant, chronic renal disease, asplenia), see resources in footnote b.
Can patients with	Also see separate section on immunosuppressive MEDICATIONS, below
immunocompromise receive LIVE vaccines?	General concepts: Avoid live vaccines unless immunocompromise is mild, data supports use of the vaccine, and the risk of natural infection is greater than the risk of immunization. ³ Live vaccines should not be given to severely immunocompromised patients, or if immune status is uncertain. ^{1,3} The ultimate determination of severe immunocompromise should be made by the provider treating the patient's immunocompromising condition. ^{1,3}
	Special disease-considerations (medications are discussed below):
	 Some patients with B-cell deficiency can receive certain live vaccines.^{1,3} For details, see resources in footnote b. Live vaccines are not contraindicated in patients with complement deficiency.^{2,3} HCT: Live vaccines should not be given within four weeks of the onset of the pre-transplant conditioning regimen.³ BCG should never be given to any patient who might need an HCT.³ MMR and varicella vaccines can be given to HCT recipients 24 months post-transplant, assuming immunocompetence.¹ Solid organ transplant: live vaccines should be given at least four weeks prior to transplant.³ Live vaccines are generally contraindicated post-transplant.³ Asplenia: only LAIV (e.g., FluMist) is contraindicated (U.S.).² HIV patients who are not severely immunocompromised can get MMR, varicella, and rotavirus.^{2,3} For help identifying these patients, see resources in footnote b.

Clinical Question	Pertinent Information of Resource
Can patients receiving	General concepts
immunosuppressive MEDICATIONS	• Because inactivated vaccines cannot replicate, they are safe for immunocompromised patients. However, these patients may not respond as well as immunocompetent patients. Consider the following:
receive vaccines?	o Review vaccination history and administer any needed inactivated vaccines at least two weeks before planned immunosuppressive therapy to optimize response.³ In addition to vaccines recommended as for immunocompetent patients, other vaccines may be recommended: ■ Pneumococcal vaccination. ^{2,3,19} For guidance, see reference 11 (Canada) or 12 (US). ■ Recombinant zoster vaccine (Shingrix) is recommended for adults ≥19 years of age (US). ^{2,19} ■ Meningococcal vaccination is recommended for patients who will receive eculizumab (Solaris) or ravulizumab (Ultomiris). ² For guidance, see references 11 (Canada) or 12 (US). O For patients already on immunosuppressive therapy: ■ Response varies depending on the vaccine, drug, and patient population, and is generally attenuated; nevertheless, patients receiving immunosuppressive therapy can benefit from vaccination. ^{5,9,16,17} ■ If risk of infectious exposure is low, consider delaying inactivated vaccines until the person is less immunosuppressed. ³ ■ In adults receiving immunosuppressants for rheumatic disease, consider using a high-dose or adjuvanted influenza vaccine, if available, instead of a standard influenza vaccine. ¹⁹ ■ If the patient is vaccinated during immunosuppression, consider checking titers once the drug is discontinued to guide whether vaccination requires repeating. ³ ■ In chemo patients, expect vaccines to be held during chemo, although an inactivated vaccine (e.g., influenza) might be given between cycles. Patients might be revaccinated with vaccines given during chemo when chemo is over. ² Each center will have protocols. ■ Canada: double the usual hepatitis B vaccine dose, and use a 3- or 4-dose schedule. ³ ■ HPV vaccine (e.g., Gardasil 9) should be given using a 3-dose schedule ³ . ¹¹² ■ In general, live vaccines should be avoided in patients receiving high-level immunosuppressive therapy to reduce risk of acquiring an infection from the vaccine. ^{2,3}
	Varicella vaccination is recommended for susceptible patients before IBD immunosuppressive therapy is started.8
	 Specific medications: Consult prescribing information/product monographs for MS therapies for guidance. Also see below concerning use of alemtuzumab for cancer. Deucravacitinib for psoriasis: discontinue two to three half-lives prior to live vaccination, and restart two to four weeks post-vaccination.¹⁸ Non-live vaccines can be given without deucravacitinib interruption.¹⁸
Continued	• Cyclosporine for psoriasis: hold for two to four weeks after live vaccination. Non-live vaccines can be given without cyclosporine interruption. 18

Clinical Question	Pertinent Information of Resource
Can patients receiving	Tofacitinib for rheumatic disease or psoriasis: hold for one week prior to live vaccination, and restart two to four
immunosuppressive	weeks after live vaccination (ACR: hold for four weeks post-vaccination). Non-live vaccines can be given without
MEDICATIONS	tofacitinib interruption. 18,19
receive vaccinations, continued	• Leflunamide, mycophenolate, calcineurin inhibitors (e.g., cyclosporine), or oral cyclophosphamide for rheumatic disease: hold for four weeks prior to live vaccination, and restart four weeks after live vaccination. Non-live vaccines can be given without treatment interruption.
	• Low-level immunosuppression (see footnote a): varicella can be given. ³ Other live vaccines can be given after a
	risk/benefit assessment (e.g., MMR before travel). ^{3,5} Consult an expert if immunosuppressants are used in combination. ³
	Methotrexate for rheumatic disease or psoriasis: consider holding for two to four weeks prior to live vaccination, and restarting two to four weeks after live vaccination (ACR: hold methotrexate for four weeks before and after live vaccination. Hold times can be shorter if live vaccination is critical and disease flare risk is high.). 18,19
	Consider holding methotrexate for two weeks after non-live vaccines (including COVID-19), if disease activity allows. ^{18,20} (ACR: consider holding methotrexate for two weeks after non-live influenza vaccine, if disease
	activity allows, but other non-live vaccines can be given without methotrexate interruption [COVID-19 not addressed]. ¹⁹)
	O Azathioprine for rheumatic disease : hold for four weeks prior to live vaccination, and restart four weeks after live vaccination. Hold times can be shorter if live vaccination is critical and disease flare risk is high. Non-live vaccines can be given without azathioprine interruption.
	• High-level immunosuppression (see footnote a): IBD guidelines recommend a three-month washout of
	immunosuppressive therapy before giving live vaccines (four months for the yellow fever vaccine).8
	O Biologics : live vaccines should be avoided in patients receiving biologics (e.g., therapeutic monoclonal antibodies, [e.g., adalimumab, etanercept, infliximab, etc], lymphocyte-depleting agents).
	 Some rheumatologic experts recommend a washout of two to three half-lives before giving live vaccines (at
	least four weeks) and restarting two to three half-lives after administration of live vaccines (at least one to two
	weeks). 14 IBD guidelines recommend a three-month washout from high-level immunosuppressive therapy (see footnote a) (four months for the yellow fever vaccine). 8
	• Rituximab or alemtuzumab may cause prolonged immunosuppression. ¹ Some experts advise waiting at least six to 12 months after treatment to vaccinate. ^{3,5} B cell enumeration is generally performed during rituximab therapy and should be reviewed prior to immunization. ³ Although data is lacking, some experts would recommend waiting at least four weeks after vaccination to restart rituximab. ^{1,5}
	o Rituximab for rheumatic disease: consider giving non-live influenza vaccine when appropriate, but
	consider deferring other non-live vaccines until the next rituximab dose is due. Wait two weeks post- non -
	live vaccination to restart rituximab, if disease activity allows. 19 TNF inhibitors, IL-12/IL-23 inhibitors, IL-23 inhibitors, IL-17 inhibitors, IL-6 inhibitors, IL-1 inhibitors,
	or belimumab for psoriasis or rheumatic disease: discontinue two to three half-lives prior to live vaccination,
	or beamanable poortages of rheumane disease. discontinue two to three hair fives prior to five vaccination,

Clinical Question	Pertinent Information of Resource
	and restart two to four weeks post-vaccination (ACR: hold for one dosing interval prior to live vaccination, and restart four weeks post-vaccination). Non-live vaccines can be given without treatment interruption. Anifrolumab for rheumatic disease: hold for one dosing interval prior to live vaccination, and restart four weeks post-vaccination. Non-live vaccines can be given without anifrolumab interruption. Abatacept for rheumatic disease or psoriasis: discontinue four weeks (intravenous) or one week (subcutaneous) prior to live vaccination, and restart two to four weeks post-vaccination (ACR: hold for one dosing interval prior to live vaccination, and restart four weeks post-vaccination. Non-live vaccines can be given without abatacept interruption. 18,19 Non-live vaccines can be given without abatacept interruption. 18,19 Non-live vaccines can be given without abatacept interruption. Non-live vaccines can be given without cyclophosphamide interruption. and restart four weeks post-vaccination. Non-live vaccines can be given without cyclophosphamide interruption. If a cancer patient is at least three months post-chemo/radiation, and restart four weeks post-vaccines can be given. Rituximab and alemtuzumab are exceptions (see above). In Immunosuppressive corticosteroid dose (see footnote a): Live vaccines should be deferred for at least four weeks after stopping an immunosuppressive corticosteroid dose. It live vaccines should be deferred for at least four weeks after stopping an immunosuppressive corticosteroid dose. Was guidelines recommend a three-month washout after high-dose, systemic corticosteroids taken for ≥2 weeks, or one month after a short-term, high-dose pulse. Wait four weeks post-vaccination to restart. Consider giving non-live influenza vaccine when appropriate, but consider deferring other non-live vaccines until the corticosteroid dose can be tapered to the equivalent of prednisone <20 mg/day.
Can HOUSEHOLD CONTACTS of immunocompromised patients receive LIVE vaccines?	 Household contacts may receive MMR, varicella, rotavirus, and LAIV (e.g., FluMist).^{1,3} See resources in footnote b for other vaccines recommended for contacts. If a recipient of the varicella vaccine develops a rash, they should keep the rash covered and avoid direct contact with the immunocompromised person until the rash has cleared.^{3,5} LAIV (e.g., FluMist) is contraindicated in close contacts and caregivers of severely immunocompromised patients (e.g., HCT recipients requiring hospital isolation).^{3,13} Healthcare workers and visitors who have received LAIV should avoid contact with severely immunocompromised patients for seven days after vaccination (Canda: two weeks).^{3,13} Immunocompromised patients should avoid handling diapers of infants within the first month of infant rotavirus vaccination.⁵

Abbreviations: ACR = American College of Rheumatology; BCG = bacilli Calmette-Guerin; HCT = hematopoietic cell transplant; HPV = human papilloma virus; Hib = *Haemophilus influenzae* type b; IBD = inflammatory bowel disease; IL = interleukin; LAIV = live attenuated influenza virus; MMR = measles, mumps, rubella; MS = multiple sclerosis; TNF = tumor necrosis factor

- a. Immunosuppressive steroid dose (i.e., high-level immunosuppression dose): prednisone ≥20 mg daily or ≥2 mg/kg daily (or equivalent) for ≥14 days.^{1,3} This does NOT include alternate-day regimen; rapid tapers; short (<14 day) high-dose regimen; topicals; physiologic replacement doses; or intra-articular, bursal, or tendon injection.¹⁻³ Live vaccines can be given to patients receiving inhaled corticosteroids (Canada: with the exception of LAIV, which should not be given to patients with severe asthma receiving high-dose inhaled corticosteroids).^{1,3}
 - **Low-level immunosuppression** examples: methotrexate \leq 0.4 mg/kg/week, azathioprine \leq 3 mg/kg/day, or 6-mercaptopurine \leq 1.5 mg/kg/day).³
 - **High-level immunosuppression** examples: immunosuppressive corticosteroid dose (see above), methotrexate >0.4 mg/kg/week, azathioprine >3 mg/kg/day, or 6-mercaptopurine >1.5 mg/kg/day; adalimumab, certolizumab, etanercept, golimumab, infliximab, natalizumab, vedolizumab. Consult prescribing information for MS treatments (e.g., fingolimod).
- b. Additional resources:
 - US: Altered immunocompetence. General best practice guidelines for immunization: best practices guidance of the Advisory Committee on Immunization Practices (ACIP). (https://www.cdc.gov/vaccines/hcp/acip-recs/general-recs/immunocompetence.html).
 - US: CDC Recommended Adult Immunization Schedule (https://www.cdc.gov/vaccines/schedules/downloads/adult/adult-combined-schedule.pdf).
 - Canada: Canadian Immunization Guide, Immunization of Immunocompromised Persons (https://www.canada.ca/en/public-health/services/publications/healthy-living/canadian-immunization-guide-part-3-vaccination-specific-populations/page-8-immunization-immunocompromised-persons.html#t5).
- c. If the drug has more than one approved dosing frequency, hold for the longest approved dosing interval; however, for IL-6 or IL-1 inhibitors, in children with systemic juvenile rheumatoid arthritis or other autoinflammatory disorder, shorter hold times can be considered if live vaccination is critical and the risk of disease flare is high.¹⁹

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