Consent Policy Design Group

Guiding Principle Compilation of Responses from Design Group Members & Suggested Recommendations September 24, 2019

Principles	Accept	Accept with Changes	Don't Accept	Unable to Respond	Proposed Changes		
 Consent policies should require patients be provided clear and unambiguous information about health information sharing choices under applicable State 	2	2			(PC) And should conform with the most recent TEFCA guidelines (policy) including the concept of		Deleted: Connecticut and Federal statutes
and Federal law. The information should be translated for non-English speakers and should conform to the Web Content Accessibility Guidelines ¹ developed by the Web Accessibility Initiative (WAI), part of the World Wide Web Consortium (W3C). ²	SB, NS	RR, PC			meaningful choice		Commented [R1]: These 2 sentences are 2 totally different
Recommendation: Accept with changes. Consent policies should require patients be provided clear and a	unambigu	ous inform	ation abou	t health info	rmation sharing choices under		concepts and requirements and should be divided into 2 different principles.
applicable State and Federal law. Other considerations:							Commented [CR2R1]: Agreement has been to move this language to a different principle. The group will need to decide which principle this should be added to.
						,	Commented [MM3R1]: Moved to GP2
 Consent policies should require Connecticut's Office of Health Strategy to develop an educational resource tool kit on health information sharing, 	3			1	(RR) See comments. (PC) Shouldn't be dependent on funding. IAPD funding should		Deleted: Dependent upon funding availability

¹ <u>https://www.w3.org/WAI/standards-guidelines/wcag/</u>

² https://www.w3.org/WAI/

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leveraging and adapting content from recognized	SB,			RR	include funds to cover this
third-party resources, ³ Educational content should be	NS, PC				component
reviewed and approved by the Health IT Advisory					
Council, and should not only include information for					
patients, parents and guardians, but also for					
providers, pharmacies, labs, health plans, state and					
local government agencies, and employers.					
Recommendation: Accept with changes.					
Consent policies should require Connecticut's Office of Health Si sharing, leveraging and adapting content from recognized third Health IT Advisory Council, and should not only include informa health plans, state and local government agencies, and employ conform to the Web Content Accessibility Guidelines ⁵ developed Consortium (W3C). ⁶	l-party res tion for po ers. The ir	sources.⁴ Ec atients, par nformation	lucational ents and g should be	content sho uardians, bi translated f	uld be reviewed and approved by the it also for providers, pharmacies, labs, or non-English speakers and should

Other considerations:

3. Information a	nd educational resources on consent			Internal Review: I can see the value
policies shoul	d be distributed broadly by <mark>Health</mark>	3	1	in having HIA disseminate this, but
Information A	lliance, Inc. (HIA) throughout			 not as the sole source of this info.

⁴ Adapted, with permission, from the CARIN Alliance Trust Framework and Code of Conduct (<u>https://www.carinalliance.com/our-work/trust-framework-and-code-of-conduct/</u>

⁵ https://www.w3.org/WAI/standards-guidelines/wcag/ ⁶ https://www.w3.org/WAI/ Deleted: to the extent possible

Commented [R6]: I have no idea who this organization is and how it relates to State policy. Why would we as the State rely on distributed resources of a private organization, and in particular of one organization over another? Shouldn't it be the State that determines our educational resources to support the State's official HIE and the functions the State decides are appropriate to run through the HIE?

³ Adapted, with permission, from the CARIN Alliance Trust Framework and Code of Conduct (<u>https://www.carinalliance.com/our-work/trust-framework-and-code-of-conduct/</u>

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Connecticut and be made widely available and easily accessible through a variety of sources including the Health Information Alliance, all health and human services agencies and departments in the state of Connecticut, and organizations participating in HIE services in Connecticut.	SB, NS, PC		RR		People aren't initially going to go there, or even know they should, to look for this. OHS should be the lead in working with stakeholders to develop a comprehensive and sustainable plan for managing consent education.

Recommendation: Accept with changes.

Information and educational resources on consent policies should be distributed broadly throughout Connecticut and be made widely available and easily accessible through a variety of sources including the Health Information Alliance, all health and human services agencies and departments in the state of Connecticut, and organizations participating in HIE services in Connecticut. The distribution process will be supported by HIA's partners, including the Office of Health Strategy (OHS).

Other considerations:

4.	A review of consent policy considerations should be conducted by HIA, Inc. for each HIE use case before	2	1	1	(PC) Refer to #1
	an HIE use case is put into production, with a use	SB, NS	PC	RR	
	case-specific consent policy developed if indicated				
	from the review.				

Recommendation: Accept with changes.

A review of consent policy considerations should be conducted for each HIE use case before an HIE use case is put into production, with a use case-specific consent policy developed if indicated from the review.

Other considerations:

Commented [R7]: Why? And would this review need to be funded? Wouldn't that need to go out to bid to then determine if in fact this is the right organization to do this review?

Commented [R8]: Why wouldn't this be reviewed by an internal State committee, similar to this Consent Design Group, filled with Subject Matter Experts and including appropriate Legal Counsel to review compliance with applicable laws and best practices in both privacy, patient choice, patient experience and quality?

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5.	participation in electronic health information exchange should be included in the Notices of Privacy Practices (NPP), as required of healthcare organizations by Health Insurance Portability and Accountability Act (HIPAA). This inclusion in the NPP	3		1		
	 should be standard practice across the state of Connecticut, whether the exchange of health data is facilitated by: a. a national consortium; b. an association of healthcare providers or hospitals on behalf of their members; c. a group of healthcare organizations operating under single tax ID for healthcare payment 	SB, NS, PC		RR		
	 under an accountable care arrangement; a group of healthcare organizations using the same electronic health record system vendor; or entities incorporated or designated for the purpose of facilitation electronic purpose of 					
	purpose of facilitating electronic exchange of health data. mendation: TBD 10/15					
other						
6.	Consent policies should result in the lowest possible burden on providers responsible for their implementation and maintenance.	2 SB, NS	1 PC		1 RR	(PC) But provide ample information for patient understanding and questions.

Commented [R9]: It is higly unlikely large organizations that participate in numerous HIE's and similar exchanges will be able and/or willing to modify their NPP's each time a new use case or exchange process is added. Healthcare Organizations should instead be required to at a minimum state in the NPP's that the organization participates in HIE's and similar exchanges, and have a location (e.g., a designated website) that could list the various HIE's and exchanges they participate in over time, with links to further explanations and guidance from the entity, OHS's resources and other information about exchanges and who to contact with questions or concerns. Including this specific requirement could result in certain organizations deciding not to participate in this specific HIE, particularly where other HIE's are available that would eventually connect to this HIE in all likelihood anyway.

Commented [R10]: This is a great statement in theory, but depending upon the size and perspective of the provider, I'm not sure we can define what constitute a burden for them.

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Recommendation: TBD 10/15							
Other considerations:							
7. Clearly written information about consent policy					Internal Review: Seems closely	1	
changes should be provided by	3			1	related to #3. Maybe consolidate		Commented [R11]: Again, why would this be provided by
to patients, parents and guardians, state and local					this message?		HIA rather than OHS?
health and human service agencies, and all licensed	SB,			RR			Deleted: HIA, Inc
healthcare entities in a timely manner when policies	NS, PC						Deleted:
or practices have changed, adhering to the principles							
of broad dissemination and accessibility of							
information described above. ³ Recommendation: TBD 10/15						-	
Other considerations:							Commented [R12]: Yes, I agree. This should go without saying. But without clarity as to how this would be accomplished and what it might look like from the Privacy- By-Design perspective, it is hard to say whether what mechanisms would be appropriate. It is also not clear from
8. Mechanisms, including digital tools, for expressing	3			1	(PC) Implies everyone has access to		how this is worded what "digital tools" would mean. Does that mean there is a way for an individual patient to login to
consent policy preferences should be user-friendly	SB,			RR	and basic literacy to use digital		the HIE and consent? Is this a datafeed that happens on the
and easily accessible.	NS, PC				tools. Should there be a paper	J	Healthcare Organization's side that blocks a patient's
					option		information from going to the HIE is affirmative consent is not received? Or if we are going via an opt-out
Recommendation: TBD 10/15 Other considerations:							methodology (and again, we have not determined as group whether a recommendation for opt-out vs. a recommendation for opt-in is more appropriate for consent
		1		1	1		for this HIE or other exchanges using State tools), how that would be addressed digitally or electronically? We do not have enough information as a group to even say whether
9. Consent policies should explain what will happen to	5				(PC) Implies a mechanism to		this very basic thing that we probably could all agree to as a
the patient's data and previously provided	SB,				withdraw consent not previously		group as a recommendation is even appropriate.
authorization, if consent is withdrawn.	NS,	<u> </u>		1	noted.		Formatted: Tab stops: 1.44", Left

Deleted: after they withdraw their consent

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	nd previous	ly provided	authorizati	on, if consent is withdrawn.	-	
4			1	(PC) Does this suggest they may exchange information outside the bounds of the acted use cases or	-	Commented [R13]: This is not a Consent policy requirement; this is a HIPAA Privacy regulatory requirement. Why would this even go in a consent policy?
SB, NS, SI, PC			RR	HIPAA allowed uses? Need more info to sign off.		
s. deletion	<u>l.</u>		1			
3						Commented [R14]: Again, these are HIPAA Privacy Rule,
	A RR, PC, SI t's data ar 4 SB, NS, SI, PC	Accept with Changes RR, PC, SI t's data and previous 4 SB, NS, SI,	Accept with Changes Don't Accept RR, PC, SI PC t's data and previously provided 4 SB, NS, SI, PC PC	Accept with Changes Don't Accept to Respond RR, PC, SI PC, SI Image: Signal state	Accept with Changes Don't Accept to Respond RR, PC, SI Proposed Changes t's data and previously provided authorization, if consent is withdrawn. t's data and previously provided authorization, if consent is withdrawn. 4 1 SB, NS, SI, PC RR NS, SI, PC RR	Acceptwith ChangesDon't Acceptto RespondProposed ChangesRR, PC, SIPC, SIImage: State of the stat

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12. Consent policies should address sensitive and specially protected data, including, but not limited to, mental health, substance abuse, and HIV status data, in alignment with federal and state statutes. Recommendation: Accept with changes.	3 SB, NS, PC			1 RR	(PC) This should be a shall	 Commented [R15]: Again, in theory I agree with this concept, but we would need clarity on the use cases where specially protected data would pass through the HIE in the first place. Many of the regulations involving these specially protected data categories specifically require affirmative concert, is a sub-transformative special of the regulation of the regulations in the second of the regulations in the second of t
Consent policies shall address sensitive and specially protected sexually transmitted disease data, as well as age-related privat		-				consent (i.e., authorization, explicit opt-in, etc.) to share data. This would be again then be a regulatory requirement, not a consent policy nicety.
13. Consent policies should be aligned with certain national interoperability initiatives, including the Common Agreement (CA) under development as part of Trusted Exchange Framework and Common	3 SB, NS, PC			1 RR	-	 Commented [R16]: It is hard to comment on this proposed principle without clarity as to what is meant by the term "be aligned with"
Agreement (TEFCA), to support the ability to exchange data with entities outside the state. Recommendation: TBD 10/15	N3, PC					
Other considerations: 14. Consent policies should be reviewed annually (or	3			1		 Commented [R17]: Is annually practical? A specific consent
biannually) to ensure it is aligned with these principles and complies with any changes in best practices or federal or state law.	SB, NS, PC			RR		process should be reviewed each time a new use case is to be implemented. The over-arching policy should also be reviewed when new use-cases are contemplated to make sure everything still lines up. But I think saying annually or
Recommendation: Accept with changes. Consent policies should be reviewed periodically to ensure it is federal or state law. Other considerations:	aligned w	ith these pr	inciples and	d complies v	with any changes in best practices or	biannually, as opposed to something like, as a minimum when new use cases or regulatory changes may impact consent, provides more flexibility and is probably more manageable. Also, who would do this review? Would it be OHS? A consent process committee?

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 Consent policies should provide a clear procedure for addressing complaints by individuals regarding the use of their data. 	5 SB, NS, RR, PC, SI				(PC) Should thid be a special Board or Committee? Internal or external?
Recommendation: Accept as written.					
Other considerations:					
16. OHS should consider pursuing regulations that define requirements for compliance with consent	3 SB,			1 RR	-
policies.	NS, PC				
Recommendation: TBD 10/15 Other considerations:					
17. Consent policies should require that patients have	3			1	
sufficient time to review educational material before making a consent decision. ⁷	SB, NS, PC			RR	
Recommendation: TBD 10/15					
Other considerations:					
	5				

⁷ Adapted from ONC, HealthIT.gov Meaningful Consent Overview (<u>https://www.healthit.gov/topic/meaningful-consent-overview</u>)

Commented [R18]: What does this mean? Why would there be a time crunch for a patient to decide? Again, whether we do an opt-in or an opt-out may matter here. If sharing is not going to be automatic, then a patient should be able to take as long as they like to decide. If the process is going to be automatic based upon a decision to put this simply in NPP's and then let patients opt out either affirmatively, or by choosing not to receive healthcare at an organization that participate in this and/or exchanges, then this time element becomes irrelevant.

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18. Consent policies should require a consent decision is	SB,				
not used for discriminatory purposes or as condition	NS,				
for receiving medical treatment ⁴ .	RR,				
	PC, SI				
Recommendation: Accept as written.					
Other considerations:					
19.					
20.					
21.					
22.					
23.					
24.					
25.					

In Reference to Guiding Principle #1:

Susan Israel - The public needs to be informed whether CT is an opt-out state for transparency and in order for the "unambiguous information about health sharing choices," name the groups (who) will be seeing their data (identified or unidentified) and where the data would be sent and stored temporarily or permantently. It must be stated for which data the patients can exercise consent vs what are automatically sent by their providers by virtue of recievnig medical care:

- 1) identify and care mapping data
- 2) identity quality control
- 3) clinical care summaries with diagnoses and medications
- 4) empanelment
- 5) quality data measurement output
- 6) public health beyond communicable diseases

- 7) opioid medication that includes psychiatric drugs which are also controlled substances
- 8) outpatient and inpatient discharge summaries
- 9) identified tumor registry
- 10) lab and imaging data
- 11) individual electronic health record longitutidal data
- 12) Surecripts data (from which there is an opt-out?)

Obviously, there is the tension of patient rights to opt-out of an electronic health infomation exchange for their mandated data to go to OHS/CT. But the removal of patient right of consent for the sharing of their data for TPO which includes oversight was removed by HHS in 2003 without Congressional approval. CT taking so much private medical data (including the APCD) and the HIPAA rules certainly stretch what the government is "allowed" to do under the 4th amendment.