Community Affairs Legislation Committee
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RE: APF submission; Inquiry into Healthcare Identifiers Bill 2010 and Healthcare Identifiers (Consequential Amendments) Bill 2010

The Australian Privacy Foundation (APF) has been active since 1987 as the country's leading public interest advocacy organisation focussing specifically on privacy. I am writing in my capacity as Chair of the Health Sub Committee of the APF.

Executive Summary
The APF has made many attempts to communicate with NEHTA and the Department of Health and Ageing on the succession of eHealth initiatives over the last decade. Despite claims to the contrary, effective consultation with consumer advocacy NGOs has emphatically not yet taken place. Key points made in the APF submission are summarised below.

1. The Health Identifier (HI) Bills contradict the APF Policy Statement on eHealth Data and HIs. It is attached, and at http://www.privacy.org.au/Papers/eHealth-Policy-090828.pdf.

Accordingly the APF opposes the Healthcare Identifiers Bill and the Healthcare Identifiers (Consequential Amendments) Bill 2010 entirely.

2. Even if a HI were to be included in a comprehensive eHealth scheme, the APF opposes the piecemeal approach the Government has adopted. The current Bills provide only a tiny fraction of a complete plan. They cannot be understood in the absence of enabling legislation for information flows, authentication, reliability and health information privacy aspects.

In any case, the APF draws to attention the impossibility of evaluating the utility of the HI system for patient privacy and health when only a fraction of the proposal is on the table, and even the relevant agencies appear to know little about how it would work in a “real-life” context.
3. Should the HI be authorised by the Parliament in the absence of a comprehensive eHealth plan, these Bills are seriously deficient in relation to such matters as:

1. the absence of a coherent and convenient mechanism whereby individuals will know what their own HI is;
2. the absence of information relating to access by consumers or their carers to patient data that becomes linked to the HI;
3. the impossibility of providing access control over a scheme to which more than half-a-million individuals would have access;
4. the ineffectiveness of access logging, and hence the unenforceability of penalties on individuals who commit or enable information security breaches;
5. the failure to impose penalties on organisations that commit or enable information security breaches, including servants of the Crown, and the failure to create an enforcement regime to ensure organisations comply; and

In any case, the APF draws attention to serious deficiencies in the Bill itself.
APF submission; Inquiry into Healthcare Identifiers Bill 2010 and Healthcare Identifiers (Consequential Amendments) Bill 2010

Introduction

The Australian Privacy Foundation is the primary non-governmental organisation dedicated to protecting the privacy rights of Australians. The Foundation aims to focus public attention on emerging issues which pose a threat to the freedom and privacy of Australians. Since 1987, the Foundation has led the defence of the right of individuals to control their personal information and to be free of excessive intrusions. For information about the Foundation see www.privacy.org.au.

Background

The APF has made many attempts to communicate with NEHTA and the Department of Health and Ageing on the succession of eHealth initiatives over the last decade. However, both agencies have avoided engagement with privacy advocates. The few events held have been sporadic and have involved the agencies talking, but not listening, to advocates. Contrary to claims by the agencies, effective consultation (i.e. meaningful two way communication) with consumer advocacy NGOs has emphatically not taken place.

APF Position

The APF issued its Policy Statement on eHealth Data and Health Identifiers in August 2009. It is attached and is also published at http://www.privacy.org.au/Papers/eHealth-Policy-090828.pdf. Relevant aspects of the policy are as follows:

1. The health care sector must remain a federation of islands
2. Consolidated health records must be the exception not the norm
3. Health identifiers must be at the level of individual applications
4. Pseudo-identifiers must be widely-used
5. Anonymity and persistent pseudonyms must be actively supported.

Accordingly the APF opposes the Healthcare Identifiers Bill and the Healthcare Identifiers (Consequential Amendments) Bill 2010 entirely.

Deficiencies in the Bill

If, despite the inappropriateness of the policy, and despite the tiny proportion of the complete package of legislation that is necessary in order to understand the proposal, the Parliament considers the Healthcare Identifiers Bill 2010 and Healthcare Identifiers (Consequential Amendments) Bill 2010 in isolation, then they would need to be substantially amended to overcome specific deficiencies.

The APF submission addresses the issues outlined for the Committee to consider during this enquiry. The comments below are therefore mainly confined to the following matters:

1. Operation of the Healthcare Identifier Service, including access to the Identifier
2. Privacy safeguards in the Bill
3. Relationship to national e-health agenda and electronic health records.

We have also attached a copy of our submission to with regard to the exposure draft Healthcare Identifiers Bill 2010 of January 6 2010 for background information.

1. Operation of the Healthcare Identifier Service, including access to the Identifier

Figures released by the National E-Health Transition Authority (NEHTA) indicate that more than 500,000 Healthcare Provider Individuals (HPIs) will be able to use and disclose the HIs to provide patient care as is indicated in Sections 18 (Disclosure to healthcare recipient) and 23 (Disclosure to healthcare recipient) of the HI Bill. The latter does not mention patient or consumer access to the personal information linked to the HI [1, p.6].

Direct consumer access to the HI and personal information linked to it should be integrated into the HI Bill currently before parliament.

The Bill suggests personal information stored about individuals that is linked to the HI will not be directly and immediately available to the people it concerns, neither will the HI itself, except via a third party. To ensure trust in the system, patients and consumers will need to believe the 500,000 HPI workers will not make a single human error while using and disclosing their HI nor will they use the number to obtain personal information, such as addresses or names for criminal purposes or just for curiosity [2]. However records published in Personal Information Digests published by the Office of the Privacy Commissioner indicate the reverse. In 2007, 2008 and 2009, many of Medicare’s 6,000 employees, 16% of them, are suspected of using patient files for non-Medicare related purposes [3]. When one extrapolates this figure, the honey pot posed by storing the birth to grave personal information of all Australians implies that perhaps 80,000 HPI workers may misuse patient files if the HI Bill, as drafted, is enacted by the government.

Human error that results from disseminating the HI to 500,000 Australians who will routinely use and disclose the HI will risk patient health.

Patients and consumers will be required to verify information linked to an HI when visiting a health service or perhaps a Medicare office to ensure it is reliable. The verification will occur in public locations, such as the Reception areas of health services, the triage Nurse at admission to Accident and Emergency or perhaps, in a Medicare retail outlet. Patient and consumer stress reactions apparently escalate when presented with private information in public environments [4]. Anecdotal evidence and media reports suggest many patients and consumers turn to the health black market (e.g. Viagra, cognitive enhancement drugs such as ADD and ADHD, breast milk and testosterone) and self-diagnose rather than confirm personal details in public environments. The ostensible government “ownership” of personal information is vexatious to these individuals, many of whom are unlikely to verify that the information stored on their HI is accurate and to confirm details linked to the number with administrative staff in public location.

The lack of direct consumer and patient access to their HI is likely to swell the health black market as individuals self diagnose to protect their privacy.
A newspaper recently reported that many consumers will only discover they have been assigned an HI when they seek health care [5]. Health authorities will use the HI to index and share existing medical records from July 1, 2010. Details linked to the card will be considered reliable for medical treatment purposes from then. These details should be verified before treatment, but this will not necessarily be the case. For instance many patients are unconscious and unaccompanied when admitted to hospital. The information may also have been amended as a consequence of identity fraud or human error. Such mistake may well result in medical error – ‘the wrong drug’ for the ‘wrong person’ at the ‘wrong time’. Consumers and patients must be personally informed of the HI and links to their Medicare card so they are regularly empowered to check the veracity of information linked to their identity number.

### Australian consumers and patients must be personally informed of the HI service and links to their Medicare card to avert potential medical error.

The HI Bill makes no reference to the arrangement for consumer access to their HI or the attached records. The HI (Consequential Amendments) Bill 2010 simply refers to the HI-Security and Access Framework, currently in its first version [1]. There are no timelines associated with assurances about data access to consumers and patients in either bill, nor in documentation supporting the Bills. When health officials were asked about the matter in Canberra during November last year, they explained individuals will not have any direct mechanism by which to access their HI and associated information from July 1 2010, when the HI Bill is likely to take effect.

### Patients and consumers require private and direct means of access to the HI and linked personal information from the time the HI Bill takes effect in order to verify that personal information stored about them is reliable.

#### 2. Privacy safeguards in the Bill

Safeguarding privacy in the HI Bill is ostensibly a useful precaution. Yet if consumers and patients do not have direct access to the HI at the same time as the Bill comes into effect, the safeguards become illogical. It is unlikely that Medicare personnel, receptionists from health organisations or triage Nurses at Accident and Emergency will volunteer information about a suspected data breach to the individual concerned. Data breach is simply an effective way to work in an untenable clinical context for many of health workers (6). The safeguards will often depend upon direct consumer or participant access to the HI and linked personal information.

### Direct consumer and patient access to the HI and associated information need to be explicitly addressed in the HI Bill so that the safeguards it enshrines are effective.

The HI Bills and other Bills affecting health privacy have consistently overlooked the history of breaches to patient information, which have regularly occurred over several decades [6-8]. Privacy safeguards in the Bill, as has been the case for decades, hold clinicians and other health workers liable for breach of the HI data and associated information [1]. At the same time, the HI Bill indemnifies the Crown and so its agents for data breach. Health authorities are generally responsible for supplying or funding clinical infrastructure. The underlying cause of patient data
breaches is consistently linked to poor health infrastructure in care contexts. This fact has been borne out by both Australian and international researchers. The HI Bill does not hold the right health provider personnel into account for such breaches, indemnifying the Crown, and therefore exposes patients to ongoing privacy risks in the health context.

**Safeguards to privacy in the HI Bill need to be expanded to specify penalties for the organisations or individuals that provide deficient infrastructure in patient care settings.**

Subclause 29(2) of the HI (Consequential Amendments) Bill 2010 refers to

... audits to be undertaken by the Privacy Commissioner as could be undertaken in relation to personal information.

Audits relying upon electronic logons or audit logs do not prove the identity of the individual committing such breaches. Health professionals have been documented sharing user names and passwords to enable patient eHealth care for decades, despite audits that have occurred [7, 9]. Yet this is not reflected in current, publicly available, audit reports other than as noted in the Security Review conducted by NEHTA in 2009 [13]. The audit of computer logs can only prove information about machine account “so and so” that inappropriately utilised an HI and/or related personal information. Logically, unless a camera or other technology is pointed at computerised health information systems 24/7/365 no-one may ever be able to be held accountable for data breach.

The existing audit mechanisms in the HI Bill and supporting documentation are unlikely to be effective unless the mechanisms can provide evidence that will satisfy the Australian Courts as to the identity of an individual or individuals responsible for data breach.

3. **Relationship to national e-health agenda and electronic health records.**

The relationship between the HI Bills and the national e-health agenda and electronic health records endeavours depends upon the way in which electronic health records are defined. Most health services have computers and store both personal patient information and health records on these according to their own identification number. From July 1 2010, the health services can use the HI to link all of the existing files together in a searchable format – another honey pot for would-be miscreants. Therefore claims that actual stakeholder briefings have occurred on this matter are mistaken. In every forum, health authorities directed audiences to exclude discussion of the patient and consumers health information linked to an HI during question times. Discussion of the HI and current e-health systems has been completely overlooked.

Public and stakeholder discussion on data linkage between current e-health systems and the HI need to occur before the Bill proceeds.

Health authorities have repeatedly dissociated the HI from future e-health patient care in forms and publications. At the same time, the very reason given for the existence of the Bill is purportedly to enable a future SIEHR or PHR. Such discussion has been
illogical because Australians need to know the shape of the future record before
deciding on the need to any kind of an HI.

The APF questions the purpose of devising a national identity structure to link
health records that may never be implemented while the effect of the HI Bill on
actual e-health data has been overlooked.

Finally, the HI Bill enshrines the personal consumer and client information that will
be stored by health authorities to develop the system under discussion. It is not future-
proof and is based on an outmoded technology. Researchers in both Australia and
overseas have developed identifier systems that do not require direct and open linkage
to personal information.

1. References in the HI Bill to specific data-items, such as name, date of birth
   etc., need to be removed so that as technology advances so too can Australian e-
   health systems.
2. Outmoded technology must not be enshrined in the HI Bill.

4. Miscellaneous

As pointed out time and again in written submissions from the APF to the
government, comparisons between the HI Bills, the former Australia Card and Access
Card programs are striking [10, p. 9). The HI database will be the most complete and
up-to-date Australian data base/honey pot of citizen information in existence. The informa-
tion held on the Medicare database will include the personal information of all Australians
including full name, former name, address, former addresses, age, sex and birth order.
Linkage between HI data and other data, such as that held by employers, banks or
insurers are completely unacceptable.

Consumers need to be provided with ironclad assurance that “function creep”
will not occur with regard to information linked to an HI.

The findings of the now three successive PIAs on the HI proposal have been
overlooked. While health authorities will argue the identifier proposal has changed,
many of the issues addressed in those PIAs (at significant taxpayer expense) remain
relevant. Health authorities should be required to respond to the PIA
recommendations to aid consideration of the Bill for this Inquiry [13].

Health authorities must respond to the findings of the 3 HI PIAs published by
NEHTA to aid consideration of the HI Bills.

Finally, there will be no widespread pilot of the HI system before rollout. There is no
governance, no evidence suggesting the system is reliable and safe, that there are no
bugs and we have no way to assess whether abuse of the system is possible. Neither
do we understand the designers’ assumptions, which are inherent to system
development. Consumers are advised about the benefits of the implementation
throughout publications supporting the HI Bill but not the risks (6, 9). No system is
ever completely secure and therefore, neither can information stored on the system be
secure, yet this crucial point is not made at all. The HI service has been presented to
the public in an information vacuum of all the proposed applications or uses of the service.

The APF is concerned about relying on the utility of the HI system for patient privacy and health when authorities know so little about it in a “real-life” context.

Pointing to rapidly expanding medical identity thefts in the United States in a recent edition of the MJA, the author exhorts Australian authorities to learn from international experience before the same occurs here [10]. Thus, the APF beseeches the Government to reject the HI Bills entirely. If determined to on follow the path outlined in Healthcare Identifiers Bill 2010 and Healthcare Identifiers (Consequential Amendments) Bill 2010, then we ask that the Bills be substantially amended in accordance with this submission.

Yours sincerely

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References


This document builds on the APF’s submissions over the last two decades, and particularly during the last three years, in order to consolidate APF’s policy position. It presents a concise statement of general Principles and specific Criteria to support the assessment of proposals for eHealth initiatives and eHealth regulatory measures.

The first page contains headlines only, and the subsequent pages provide further explanation.

**General Principles**

1. Health Care Must Be Universally Accessible
2. The Health Care Sector is by its Nature Dispersed
3. Personal Health Care Data is Inherently Sensitive
4. The Primary Purpose of Personal Health Care Data is Personal Health Care
5. Other Purposes of Personal Health Care Data are Secondary, or Tertiary
6. Patients Must Be Recognised as the Key Stakeholder
7. Health Information Systems are Vital to Personal Health Care
8. Health Carers Make Limited and Focussed Use of Patient Data
9. Data Consolidation is Inherently Risky
10. Privacy Impact Assessment is Essential

**Specific Criteria**

1. The Health Care Sector Must Remain a Federation of Islands
2. Consolidated Health Records Must Be the Exception not the Norm
3. Identifiers Must Be at the Level of Individual Applications
4. Pseudo-Identifiers Must Be Widely-Used
5. Anonymity and Persistent Pseudonyms Must Be Actively Supported
6. All Accesses Must Be Subject to Controls
7. All Accesses of a Sensitive Nature Must Be Monitored
8. Personal Data Access Must Be Based Primarily on Personal Consent
9. Additional Authorised Accesses Must Be Subject to Pre- and Post-Controls
10. Emergency Access Must Be Subject to Post-Controls
11. Personal Data Quality and Security Must Be Assured
12. Personal Access and Correction Rights Must Be Clear, and Facilitated
General Principles

1 Health Care Must Be Universally Accessible. Access to health care must not be conditional on access to health care data or on demonstration of the person’s status (such as residency rights or level of insurance)

2 The Health Care Sector is by its Nature Dispersed. Health care is provided by thousands of organisations and individual professionals, each with a considerable degree of self-responsibility. The sector is far too large, and far too complex to be centrally planned. Instead it must be managed as a large, complex and highly de-coupled system of autonomous entities, each of which is subject to regulation by law, Standards and Codes

3 Personal Health Care Data is Inherently Sensitive. Many individuals have serious concerns about the handling of at least some categories of health care data about themselves. Their willingness to divulge important information is important to their health care, but is dependent on them having confidence about how that information will be managed

4 The Primary Purpose of Personal Health Care Data is Personal Health Care. The protection of the individual person is the primary function of personal health care data and systems that process it. The key users of that data are health care professionals

5 Other Purposes of Personal Health Care Data are Secondary, or Tertiary. Public health is important, but is a secondary purpose. Administration, insurance, accounting, research, etc. are neither primary nor secondary but tertiary uses. The tail of health and public health administration and research must not be permitted to wag the dog of personal health care

6 Patients Must Be Recognised as the Key Stakeholder. Government agencies and corporations must directly involve people, at least through representatives of and advocates for their interests, in the analysis, design, construction, integration, testing and implementation of health information systems

7 Health Information Systems are Vital to Personal Health Care. People want systems to deliver quality of service, but also to be trustworthy, transparent and respectful of their needs and values. In the absence of trust, the quality of data collection will be greatly reduced

8 Health Carers Make Limited and Focussed Use of Patient Data. Health care professionals do not need or want access to their patients' complete health records, but rather access to small quantities of relevant information of assured quality. This requires effective but controlled interoperability among health care data systems, and effective but controlled communications among health care professionals. Calls for a general-purpose national health record are for the benefit of tertiary users (administration, insurance, accounting, research, etc.), not for the benefit of personal health care

9 Data Consolidation is Inherently Risky. Physically and even virtually centralised records create serious and unjustified risks. Services can be undermined by single points of failure; health care data isn't universally understandable but depends on context; consolidation produces a 'honey pot' that attracts break-ins and unauthorised secondary uses and creates the additional risk of identity theft; and diseconomies of scale and scope exceed economies

10 Privacy Impact Assessment is Essential. Proposals relating to personal health care data and health care information systems must be subject to PIA processes, including prior publication of information, consultation with affected people and their representatives and advocates, and publication of the outcomes of the study. Designs for systems and associated business processes must be based on the results of the PIA, and implementations must be rejected if they fail to embody the required features
Specific Criteria

1 The Health Care Sector Must Remain a Federation of Islands. The health care sector must be conceived as islands that inter-communicate, not as elements of a whole. Health care information systems must be conceived as independent services and supporting databases that inter-operate, not as part of a virtually centralised database managed by the State. Coordinating bodies must negotiate and facilitate inter-operability, not impose central schemes.

2 Consolidated Health Records Must Be the Exception not the Norm. A small proportion of the population may benefit from linkage of data from multiple sources, primarily patients with chronic and/or complex conditions. Those patients must be the subject of consent-based, specific-purpose data consolidation. This activity must not apply to people generally.

3 Identifiers Must Be at the Level of Individual Applications. Each of the large number of dispersed health care information systems must use its own identifier for people. A system-wide or national identifier might serve the needs of tertiary users of personal data, but does little for the primary purpose of personal care, and it creates unnecessary risks for individuals.

4 Pseudo-Identifiers Must Be Widely-Used. Particularly when personal data moves between organisations, the maximum practicable use must be made of one-time-use and other forms of pseudo-identifiers, in order to keep people’s identities separate from the data itself, and minimise the risk of personal health care data escaping and being abused.

5 Anonymity and Persistent Pseudonyms Must Be Actively Supported. Anonymity is vital in particular circumstances such as ensuring that people are treated for sexually transmitted diseases. Persistent pseudonyms are vital in particular circumstances such as for protected witnesses, victims of domestic violence, and celebrities and notorieties who have reason to be concerned about such threats as stalking, kidnapping and extortion.

6 All Accesses Must Be Subject to Controls. Access to personal data must be subject to controls commensurate with the circumstances, including the sensitivity of the data and the potential for access and abuse of access. This requires identification of the category of person and in many cases of the individual who accesses the data, and authentication of the category or individual identity. However, the barriers to access and the strength of authentication must balance the important value of personal privacy and effective and efficient access by health care professionals.

7 All Accesses of a Sensitive Nature Must Be Monitored. Non-routine accesses and accesses to particularly sensitive data must be detected, recorded, and subject to analysis, reporting, sanctions and enforcement.

8 Personal Data Access Must Be Based Primarily on Personal Consent. The primary basis for access to personal data is approval by the person concerned. Consent may be express or implied, and may be written, verbal or non-verbal, depending on the circumstances. All accesses based on consent must be detected, recorded and subject to analysis, reporting, investigation, sanctions and enforcement.

9 Additional Authorised Accesses Must Be Subject to Pre- and Post-Controls. All accesses that are not based on personal consent must be the subject of explicit legal authority that has been subject to prior public justification. All such accesses must be detected, recorded and subject to analysis, reporting, investigation, sanctions and enforcement.

10 Emergency Access Must Be Subject to Post-Controls. Health care professionals (but only health care professionals) must have the practical capacity to access data in apparent violation of the personal consent principle, but must only do so where they reasonably believe that it is necessary to prevent harm to some person. All such accesses must be detected, recorded, reported and subject to analysis, investigation, sanctions and enforcement.

11 Personal Data Quality and Security Must Be Assured. Data must be of a quality appropriate to its uses, and retained only as long as it remains relevant. Personal data in storage, in transit, and in use, must be subject to security controls commensurate with its sensitivity, and with the circumstances.

12 Personal Access and Correction Rights Must Be Clear, and Facilitated. Each person must have access to data about themselves, and access must be facilitated by any organisation that holds data that can be associated with them. Where appropriate, the access may be intermediated, in order to avoid misunderstandings and misinterpretation of the data. Where data is not of appropriate quality, the person must be able to achieve corrections to it.
Re: The exposure draft Healthcare Identifiers Bill 2010

The Australian Privacy Foundation (APF) is the country's leading privacy advocacy organisation. I am writing in my capacity as Chair of the Health Sub Committee of the APF.

The Foundations’ feedback to the exposure draft Healthcare Identifiers Bill 2010 is listed below.

1. The APF policy statement in relation to eHealth data and Identifiers has been brought to the attention of senior health officials and has been publicly available for several months at http://www.privacy.org.au/Papers/eHealth-Policy-090828.pdf (Appendix A). The policy, which restates submissions we have made repeatedly over many years, is completely overlooked in the draft HI Bill.

   The APF submits that the draft legislation fails to take account of significant privacy concerns despite these having repeatedly been drawn to the attention of senior health officials.

   Because this initiative is at odds with the APF’s stated policy on the matter, we reiterate our opposition to this initiative in its entirety.

   If the Department is intent on continuing down this path, despite the serious concerns, then we draw the following specific defects to your attention.

2. The draft HI Bill enables data linkage of identifiable personal information which is designed to support an electronic health record that, according to senior government officials from NEHTA and DOHA in late November last year, has not even been drafted may never be implemented.

   The purpose of devising a national identity structure to link a health record that may never be implemented is of serious concern.

3. Confusion reigns as to whether the identifiable information will be accessible to the person about whom it concerns or their carers. This is because information provided in the "Concept of Operations" and the "Update on
legislative proposals for healthcare identifiers" documents on the one hand, refer to consumer access to the HI system from implementation, but senior spokespersons from NEHTA and DoHA indicate the contrary (1,2). The spokespersons received a question directly asking about consumer access arrangements from audience members at the HI service Stakeholder Briefing Forum in the ACT on November 20 last year. Responses to the question were, firstly, that “Consumers will be given access afterwards [sic]” and secondly, that “Consumers will eventually be able to use their PIN number and a web portal. Web services will initially be given to providers but not to the consumer- this needs to be added at this stage” respectively. Nonetheless information about consumers will be available to the 600,000 health workers administering batch downloads of the HI based on Medicare or Veterans Affairs data (3). Consequently, the number will be stored in several thousand local health service information systems, regardless of their security arrangements.

The draft Bill should mandate provision for consumer access to their own HI data from the date of system implementation.

The draft Bill should mandate robust local security arrangements BEFORE authorising the storage of an HI bridge to consumers’ personal information by health services.

4. Although there may be no timelines scheduled to enable consumer access at present, several thousand publicly owned health services are expected to use and disclose the number as a unique reference number in their own health records and consumers will need to trust the security of arrangements protecting the used and disclosed information. However the data stored in the HI system may not be correct (3). Although publicly funded services will be required to use the HI from its implementation, NEHTA and DoHA spokespersons at the meeting last year suggested that in the fullness of time, consumer pressure means private practitioners will also use the HI.

We are worried by the apparent fragmentation of the health sector that is likely to occur as a consequence of silos of private sector practitioners that will not use the HI, as has been the Canadian experience (4).

5. There will be no widespread pilot of the HI system before rollout. There is no governance, no evidence suggesting the system is reliable and safe, that there are no bugs and we have no way to assess whether abuse of the system is possible. Neither do we understand the designers’ assumptions, which are inherent to system development. Consumers are advised about the benefits of the implementation throughout the “Concept of Operations” and the "Update on legislative proposals for healthcare identifiers" documents but not the risks (1, 2). No system is ever completely secure and therefore, neither can information stored on the system be secure, yet this crucial point is not made by the health authorities drafting self-referenced propaganda to support the HI system. The HI service has been presented to the public in an information vacuum of the proposed application or use of the service.

The APF is concerned about relying on the utility of the HI system for patient privacy and health when authorities know so little about it in a “real-life” context.
6. Authorities are emphasising penalties associated with the misuse and disclosure of private patient information. However research shows the overwhelming majority of misuse is actually incidental and occurs as clinicians provide patient care in environments where resources are shared (6). Disclosure occurs as people go about their ordinary work tasks— that is, no special effort is required to overhear sensitive patient information.

**Health authorities must fund a review of the context of patient care, as recently occurred in the UK, so that “incidental” privacy breaches are minimised or even eliminated completely (5).**

7. Authorities may be able to track misuse of the HI system back to the end-user account name but not necessarily to the individual concerned. Research evidence suggests the end-user account names and PKI keys are already shared with clinically unqualified and underqualified users to advance patient care tasks (6). All consumers may ever know is the account name used to obtain or disclose patient information. Also, will end-user accounts and NASH accounts be issued to individuals or to organisational units? The evidence suggests that audit information may be insufficient for consumers to use for the purposes of a complaint with regard to the misuse or disclosure of their information.

**Access control systems to support the HI seem to be inadequate with regard to protecting the privacy of Australian consumers and patients.**

8. The draft HI Bill allows data linkage between citizens’ personal information and the HI. Yet several researchers, some of whom are Australian, have developed HI systems that do not require linkages to personal information. Thus, the draft HI Bill facilitates technological approaches that are already out of date.

**The draft Bill entrenches outmoded technology into the HI system.**

9. The definitions contained in the draft Bill are open-ended and offer no guarantees to consumers at all. For example the definitions of an identified ‘healthcare provider’ and a ‘service operator’ are circular— they are whoever government authorities decide to assign a number ([page 3](#) and [page 5](#)). The use of national HI’s for "management, funding ... and the conduct of health or medical research” is also both vague and open-ended ([page 12](#)). The definition of a ‘health service’ is the most chilling of these, because during the November meeting discussed above, an insurer was advised that although they may not access the HI at this stage, **the definition of a ‘health service’ is likely to change over time.** Will HIS information be available to insurers, banks and potential employers over a period of time?

**The draft Bill would be vastly improved if it permanently closed potentially privacy invasive loopholes contained in definitions.**

10. Section 24 Regulations leaves a plethora of consumer concerns open-ended, including new data sources, governance arrangements, security mechanisms, standards, quality and safety.
The APF seeks detailed HI provisions in the legislation rather than being left to regulation on the grounds that, even though disallowable by Parliaments, the latter are rarely subject to the same level of scrutiny and debate.

11. The scope of the draft HI Bill is limited by the lack of systems to enrol some categories of Allied Health Care worker. Such enrolments may take several years to complete. A fragmented Australian HI landscape seems likely to emerge when one considers the enrolment shortcomings in the context of private health organisations, some of whom may decide not to not use an HI system, and public health organisations that will be legally required to use an HI from implementation.

Additional fragmentation due to the enrolment of Allied Health Care Workers may unintentionally exacerbate shortcomings with regard to silos of patient information (See 3, above).

12. The draft Act briefly refers to interaction with the Privacy Act in Section 6 adding yet another layer of complexity to clinical work within the Australian health privacy legal framework. Evidently, the law of a State or Territory will not apply unless it can function concurrently with the draft Bill. However nothing in the draft Bill will affect or restrict any rights or remedy a person might have had if the Bill was not enacted. The minister may revoke parts of the HI in States and Territories so long as these are advertised in the Government Gazette. The health legal landscape will be as fragmented and confusing as it has ever been, possible even more so for clinicians actually working with an HI.

Although this may be harmonised in the long term, the fragmented HI privacy legal framework outlined in the draft Bill will prove confusing for practices and clinicians, increasing the present range of threats to consumer privacy.

13. Finally, the “Building the foundation for an eHealth future …” document refers to ongoing consultations with stakeholders (1). No such consultation with most consumers or consumer groups we regularly liaise with has taken place. The overwhelming majority of consumers are excluded from NEHTA’s definition of “stakeholder” too (6). Moreover, when the APF has been invited to the so-called consultations, we have been instructed to limit feedback to the issue of HIs without reference to an EHR since at this stage we know neither whether an EHR will be enacted at all nor the nature of its relationship to an HI. However government officials can and do refer to the HIs in the context of an EHR as a way of justifying the need for HIs (2, 3). Public meetings have been similarly controlled. Consultation audiences are told what will occur, not asked about the implementation. Questions times are limited, controlled and cease well before the flood of questions do. Thus, the audience to such consultations is given little opportunity for meaningful input – the so-called consultations are little more than briefings to explain how the system will function.

APF HI policies and views are repeatedly ignored during meetings with senior health officials.
For all of the reasons herein, the APF believes the current exposure draft HI Bill is deeply flawed. It is both incomplete and inadequate in relation to privacy protection and meaningful input by health consumers. The lack of clarity around aspects of governance, slipping important issues into non existent regulations, lack of any details around NASH and lack of information about how the sector will actually use the HI service suggest to me the whole thing is under-planned and should be thought out more clearly. We are very concerned the shortcomings listed in this document will impede the development of an effective HI system for all Australians. The implication of the concerns listed above is to query whether the APF is wasting our time with this and other submissions to health care authorities.

Yours sincerely

[Signature]

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References
This document builds on the APF’s submissions over the last two decades, and particularly during the last three years, in order to consolidate APF’s policy position. It presents a concise statement of general Principles and specific Criteria to support the assessment of proposals for eHealth initiatives and eHealth regulatory measures.

The first page contains headlines only, and the subsequent pages provide further explanation.

**General Principles**

1. Health Care Must Be Universally Accessible
2. The Health Care Sector is by its Nature Dispersed
3. Personal Health Care Data is Inherently Sensitive
4. The Primary Purpose of Personal Health Care Data is Personal Health Care
5. Other Purposes of Personal Health Care Data are Secondary, or Tertiary
6. Patients Must Be Recognised as the Key Stakeholder
7. Health Information Systems are Vital to Personal Health Care
8. Health Carers Make Limited and Focussed Use of Patient Data
9. Data Consolidation is Inherently Risky
10. Privacy Impact Assessment is Essential

**Specific Criteria**

1. The Health Care Sector Must Remain a Federation of Islands
2. Consolidated Health Records Must Be the Exception not the Norm
3. Identifiers Must Be at the Level of Individual Applications
4. Pseudo-Identifiers Must Be Widely-Used
5. Anonymity and Persistent Pseudonyms Must Be Actively Supported
6. All Accesses Must Be Subject to Controls
7. All Accesses of a Sensitive Nature Must Be Monitored
8. Personal Data Access Must Be Based Primarily on Personal Consent
9. Additional Authorised Accesses Must Be Subject to Pre- and Post-Controls
10. Emergency Access Must Be Subject to Post-Controls
11. Personal Data Quality and Security Must Be Assured
12. Personal Access and Correction Rights Must Be Clear, and Facilitated
General Principles

1  **Health Care Must Be Universally Accessible.** Access to health care must not be conditional on access to health care data or on demonstration of the person’s status (such as residency rights or level of insurance)

2  **The Health Care Sector is by its Nature Dispersed.** Health care is provided by thousands of organisations and individual professionals, each with a considerable degree of self-responsibility. The sector is far too large, and far too complex to be centrally planned. Instead it must be managed as a large, complex and highly de-coupled system of autonomous entities, each of which is subject to regulation by law, Standards and Codes

3  **Personal Health Care Data is Inherently Sensitive.** Many individuals have serious concerns about the handling of at least some categories of health care data about themselves. Their willingness to divulge important information is important to their health care, but is dependent on them having confidence about how that information will be managed

4  **The Primary Purpose of Personal Health Care Data is Personal Health Care.** The protection of the individual person is the primary function of personal health care data and systems that process it. The key users of that data are health care professionals

5  **Other Purposes of Personal Health Care Data are Secondary, or Tertiary.** Public health is important, but is a secondary purpose. Administration, insurance, accounting, research, etc. are neither primary nor secondary but tertiary uses. The tail of health and public health administration and research must not be permitted to wag the dog of personal health care

6  **Patients Must Be Recognised as the Key Stakeholder.** Government agencies and corporations must directly involve people, at least through representatives of and advocates for their interests, in the analysis, design, construction, integration, testing and implementation of health information systems

7  **Health Information Systems are Vital to Personal Health Care.** People want systems to deliver quality of service, but also to be trustworthy, transparent and respectful of their needs and values. In the absence of trust, the quality of data collection will be greatly reduced

8  **Health Carers Make Limited and Focussed Use of Patient Data.** Health care professionals do not need or want access to their patients' complete health records, but rather access to small quantities of relevant information of assured quality. This requires effective but controlled interoperability among health care data systems, and effective but controlled communications among health care professionals. Calls for a general-purpose national health record are for the benefit of tertiary users (administration, insurance, accounting, research, etc.), not for the benefit of personal health care

9  **Data Consolidation is Inherently Risky.** Physically and even virtually centralised records create serious and unjustified risks. Services can be undermined by single points of failure; health care data isn't universally understandable but depends on context; consolidation produces a 'honey pot' that attracts break-ins and unauthorised secondary uses and creates the additional risk of identity theft; and diseconomies of scale and scope exceed economies

10 **Privacy Impact Assessment is Essential.** Proposals relating to personal health care data and health care information systems must be subject to PIA processes, including prior publication of information, consultation with affected people and their representatives and advocates, and publication of the outcomes of the study. Designs for systems and associated business processes must be based on the results of the PIA, and implementations must be rejected if they fail to embody the required features
Specific Criteria

1. **The Health Care Sector Must Remain a Federation of Islands.** The health care sector must be conceived as islands that inter-communicate, not as elements of a whole. Health care information systems must be conceived as independent services and supporting databases that inter-operate, not as part of a virtually centralised database managed by the State. Coordinating bodies must negotiate and facilitate inter-operability, not impose central schemes.

2. **Consolidated Health Records Must Be the Exception not the Norm.** A small proportion of the population may benefit from linkage of data from multiple sources, primarily patients with chronic and/or complex conditions. Those patients must be the subject of consent-based, specific-purpose data consolidation. This activity must not apply to people generally.

3. **Identifiers Must Be at the Level of Individual Applications.** Each of the large number of dispersed health care information systems must use its own identifier for people. A system-wide or national identifier might serve the needs of tertiary users of personal data, but does little for the primary purpose of personal care, and it creates unnecessary risks for individuals.

4. **Pseudo-Identifiers Must Be Widely-Used.** Particularly when personal data moves between organisations, the maximum practicable use must be made of one-time-use and other forms of pseudo-identifiers, in order to keep people’s identities separate from the data itself, and minimise the risk of personal health care data escaping and being abused.

5. **Anonymity and Persistent Pseudonyms Must Be Actively Supported.** Anonymity is vital in particular circumstances such as ensuring that people are treated for sexually transmitted diseases. Persistent pseudonyms are vital in particular circumstances such as for protected witnesses, victims of domestic violence, and celebrities and notorieties who have reason to be concerned about such threats as stalking, kidnapping and extortion.

6. **All Accesses Must Be Subject to Controls.** Access to personal data must be subject to controls commensurate with the circumstances, including the sensitivity of the data and the potential for access and abuse of access. This requires identification of the category of person and in many cases of the individual who accesses the data, and authentication of the category or individual identity. However, the barriers to access and the strength of authentication must balance the important value of personal privacy and effective and efficient access by health care professionals.

7. **All Accesses of a Sensitive Nature Must Be Monitored.** Non-routine accesses and accesses to particularly sensitive data must be detected, recorded, and subject to analysis, reporting, sanctions and enforcement.

8. **Personal Data Access Must Be Based Primarily on Personal Consent.** The primary basis for access to personal data is approval by the person concerned. Consent may be express or implied, and may be written, verbal or non-verbal, depending on the circumstances. All accesses based on consent must be detected, recorded and subject to analysis, reporting, investigation, sanctions and enforcement.

9. **Additional Authorised Accesses Must Be Subject to Pre- and Post-Controls.** All accesses that are not based on personal consent must be the subject of explicit legal authority that has been subject to prior public justification. All such accesses must be detected, recorded and subject to analysis, reporting, investigation, sanctions and enforcement.

10. **Emergency Access Must Be Subject to Post-Controls.** Health care professionals (but only health care professionals) must have the practical capacity to access data in apparent violation of the personal consent principle, but must only do so where they reasonably believe that it is necessary to prevent harm to some person. All such accesses must be detected, recorded, reported and subject to analysis, investigation, sanctions and enforcement.

11. **Personal Data Quality and Security Must Be Assured.** Data must be of a quality appropriate to its uses, and retained only as long as it remains relevant. Personal data in storage, in transit, and in use, must be subject to security controls commensurate with its sensitivity, and with the circumstances.

12. **Personal Access and Correction Rights Must Be Clear, and Facilitated.** Each person must have access to data about themselves, and access must be facilitated by any organisation that holds data that can be associated with them. Where appropriate, the access may be intermediated, in order to avoid misunderstandings and misinterpretation of the data. Where data is not of appropriate quality, the person must be able to achieve corrections to it.