

GDPR Policy

This Policy defines the arrangements in place within the Organisation that assures compliance to the requirements of the General Data Protection Regulation ("GDPR"), as relevant to the Organisation's business interests.

In all aspects of this Policy the term "Organisation" refers to West Valley and its services and Clinical and any attendant Services that operates as a viable and on-going business – including the employees and young people who use our services:

A: INTRODUCTION:

1. The General Data Protection Regulation (henceforth abbreviated to "GDPR") addresses certain requirements for all Organisations that collect and process personal data as part of their on-going business operations. Personal data is defined as any information relating to an "identifiable living individual", and will therefore apply to the Organisation's service users (children / young persons), employees and suppliers.

2. The GDPR applies to any data recorded in a filing system that allows personal data to be easily accessed. In this respect GDPR will apply to any of the following types of file where data may be stored:

2.1 "Hard copy" (paper) files relating to employees (e.g. employment records, safeguarding records, risk assessments, care planning, and other documents requiring original signatures).

2.2 Electronic (computer) files relating to staffing issues, (shift allocation, staff skills, training), complaints etc, and service user issues (care planning, environmental risk assessments, accidents etc).

2.3 Digital image files relating to the following:

• Photographs of staff - for ID badges / verification of identity as part of employment vetting

• photographs of young people - Care Plan / medicine assistance verification of identity / CCTV images

B: PRINCIPLES OF DATA PROTECTION:

1. The way in which our Organisation manages service user information will conform to the following 6 principles of

Information Management:

- 1. Justify the purpose(s) of using confidential information;
- 2. Only use it when absolutely necessary;
- 3. Use the minimum that is required;
- 4. Access to be on a strict need-to-know basis;
- 5. Everyone should understand his or her responsibilities;
- 6. Understand and comply with the law.



2. The Organisation is committed to the enforcement of the following Code of Good Practice in relation to the data it retains on young people and employees. In summary, data will:

- be fairly, transparently and lawfully processed;
- be used for a limited, legitimate and well-explained purpose;
- be relevant to the Organisation's needs;
- not be unnecessarily excessive in detail;

• be accurately maintained and were personal data has been identified as inaccurate, having regard to the purposes for

which they are processed, are erased or rectified without delay;

- not be kept any longer than is necessary, or as required by law;
- only be used in accordance with the individual subject's rights;

• be securely stored including protection against unauthorised or unlawful processing and against accidental loss,

destruction or damage;

• only be made available to authorised persons.

3. In this respect the following additional policies within the Organisation's documentation system are relevant:

- Confidentiality Policy
- Data Breach Policy
- C: POLICY DETAILS:

1. The core activities of the Organisation centres around the collection and processing of large quantities of personal data.

Most of this data originates from the young people as assessments of care and 'clinical' needs, care planning, records of care service delivery, risk assessments etc, but also personal data from staff members (CVs, employment records etc).

Data is classified into 2 categories for the purposes of this Policy; Personal Data and Sensitive Personal Data:

1.1 Personal Data – this is defined as any information relating to an individual who is identified or otherwise

identifiable, whether directly or indirectly. Examples here would be:

- date of birth
- current drivers' licence/passport
- photograph



• mobile phone number

1.2 Sensitive Personal Data – certain types of information have been assessed as requiring additional protection under the GDPR. This will include an individual's:

- ethnicity / racial origin
- political affiliations
- religious belief
- physical / mental health
- gender identity / sexual orientation
- genetic / biometric data (if collected)

2. This type of data requires stricter control than personal data. Staff receive appropriate training in the types of data needed to

accomplish a particular activity, for example, the development of a service user's Care Plan. Since it is highly likely that the

constituent documents and records of a Care Plan will include a mixture of both categories of data, care must be taken to

only use the data required, and not more than is absolutely necessary to deliver a high quality service.

3. The Organisation is fully committed to up-holding the legal principles of Data Protection as defined by the Office of the

Information Commissioner (ICO) In particular:

3.1 The service user will be kept informed about who, why, when, where and how ("WWWWH") their information will be held and used ("processed").

3.2 At the first point of contact, and prior to data collection, everybody whose data will be processed (current service users and staff members) will be provided with a Privacy Notice which will identify the circumstances under which any data they volunteer is collected, and subsequently processed. It is a fundamental requirement that written consent from the subject individual (service user / staff member) be obtained in order for personal data to be collected. This facility is built into the Privacy Notice.

- 4. The following will be considered when writing a Privacy Notice:
- The actual information being collected
- Who is collecting it
- How it is collected
- Why it is being collected
- How it will be used



- Who it will be shared with (as appropriate)
- The effect of this on the individuals concerned
- Whether the intended use is likely to cause individuals to object or complain

5 All individuals, young people and employees, have the right of access to manual, electronic and digital records that are relevant to their personal data. If young people are to access their residential files all 3rd Party information, information that has it has not been agreed can be shared by the author, and confidential information MUST BE removed first, if young people want to access their 'Social Work' files permission must first be gained from their social worker.).

6. Where it is deemed necessary to divulge personal data to a third party this will only be done with the express permission of the individual subject, Confidentiality Policy, In this respect both staff and service users / relatives / social workers etc will also be advised that personal information held by the Organisation may be shared with the Regulating Authority, as appropriate. However, in the interests of Safeguarding / Child Protection and when referring staff to the Disclosure and Barring Service permission will not be sought to share relevant information

The Organisation is committed to understanding and respecting the rights of the individual with respect to the safe and secure handling, storing and management of that individual's personal data. The GDPR will therefore uphold the following fundamental rights for individuals concerning their personal data:

- the right to be kept informed
- the right of access to data at any reasonable time
- the right to rectify / modify records
- the right to erase / redact any information(unless this information is known to be accurate and essential)
- the right to restrict processing of data (e.g. on a "need-to-know" basis)
- the right to data portability
- the right to object to any part of the data content

8. Personal data and records will be maintained under appropriate conditions of security to prevent any unauthorised or accidental disclosure. Records can be in hard copy (paper) format, or as electronic (word processed and scanned pdf files), or as digital files (digital photographs). In each case particular attention will be paid to the following aspects of data sharing and storage:

8.1 Hard Copy (paper) files:

- location and safety (fire, vermin) of storage;
- identification of those employees authorised to have access to specific data;
- young people / social workers authorised to have access to their personal records;
- responsibilities for secure storage of the data at the Organisation;



- retention times; i.e. how long data records are kept.
- 8.2 Electronic (computer) files:
- responsibilities for implementing data security systems for computer files;
- password-protection / encryption for access to sensitive data files;
- who is authorised to have knowledge of these passwords / encryptions;
- how often passwords are changed;
- implications for networked systems;
- how long data records are kept for (retention times);
- back-up, control and management of personal data files;
- any special control requirements needed when on-line back-up services are used.
- 8.3 Digital files (photographs, CCTV images):
- responsibilities for implementing security systems for digital files;
- password-protection / encryptions for access to sensitive data files;
- who is authorised to have knowledge of these passwords / encryptions;
- how often passwords are changed;
- how long records are kept for;
- procedures for the control and management of personal data.

9. When personal data is being processed, administrative staff will take all reasonable precautions to prevent access to data by unauthorised persons:

9.1 Record files are locked away when not required, ensuring that computer files are passwordprotected and that passwords are regularly changed.

9.2 Where practical, computer display screens are tilted towards the user and away from the general office environment.

9.3 Display screens are not left on when not in use – they should be switched off / locked.

9.4 Managing a "clear desk" policy for personal office housekeeping, ensuring that confidential electronic files are encrypted when not in use, and that waste confidential paper is destroyed by cross-cut shredding.

9.5 Ensuring that confidential conversations are not overheard.

- 9.6 Ensuring that information is transported securely.
- 10. Privacy Impact Assessments / Data Protection Impact Assessments:



Privacy Impact Assessments (PIAs), or Data Protection Impact Assessments (DPIAs), help Organizations' to identify the most effective way to comply with their data protection obligations and meet individuals' expectations of privacy.

10.1 Privacy impact assessment is a process which helps an organisation to identify and reduce the privacy risks of a project. An effective PIA uses existing project management processes to assess how a particular project or system will affect the privacy of the individuals involved.

10.2 Privacy policy, is about the right of an individual to be left alone. There are 2 types of privacy, each subject to different types of intrusion:

• Physical privacy - the ability of a person to maintain their own physical space or solitude. Intrusion can come in the form of unwelcome searches of a person's home or personal possessions, bodily searches or other interference, acts of surveillance, and the taking of biometric information.

• Informational privacy - the ability of a person to control, edit, manage and delete information about themselves and to decide how and to what extent such information is communicated to others.

Intrusion can come in the form of collection of excessive personal information, disclosure of personal information without consent and misuse of such information. It can include the collection of information through the monitoring of communications whether by post, phone or online and extends to monitoring the records of senders and recipients as well as the content of messages

10.3 Privacy Risk - the risk of harm arising through an intrusion into privacy. This is concerned primarily with minimising the risk of informational privacy - the risk of harm through use or misuse of personal information.

Some of the ways this risk can arise is through personal information being:

- inaccurate, insufficient or out of date;
- excessive or irrelevant;
- kept for too long;
- disclosed to those who the person it is about does not want to have it;
- used in ways that are unacceptable to, or unexpected by, the person it is about;
- not kept or maintained securely;
- not disposed of securely.

10.4 Harm can present itself in different ways. Sometimes it will be tangible and quantifiable, for example financial loss or losing a job. At other times it will be less defined, for example damage to personal relationships and social standing arising from disclosure of confidential or sensitive information. Sometimes harm might still be real even if it is not obvious, for example the fear of identity theft that comes from knowing that the security of information could be compromised. It can contribute to a loss of personal autonomy or dignity, or exacerbate fears of excessive surveillance.



10.5 The outcome of a PIA should be a minimisation of privacy risk. As such, the Organisation will develop an understanding of how it will approach the broad topics of privacy and privacy risks for its young people and staff.

Understanding privacy risk in this context requires an understanding of the relationship between an individual and the Organisation, and factors that can have a bearing on this include the following:

- reasonable expectations of how the activity of individuals will be monitored.
- reasonable expectations of the level of interaction between an individual and an organisation.
- the level of understanding of how and why particular decisions are made about people.

10.6 The projected benefits of a PIA - it is the Organisation's objective that undertaking an effective PIA should benefit the people affected by the project and also the organisation carrying out the project. Benefits can include the following:

• It will demonstrate to the ICO (Information Commissioner's Office) how personal data processing complies with legal requirements, and that individuals can be reassured that the organisations which use their information have followed best practice.

• A project which has been PIA assessed should be less privacy intrusive and therefore less likely to affect individuals in a negative way.

• A PIA should improve transparency and make it easier for individuals to understand how and why their information is being used.

• Undertaking the assessment will improve how the Organisation uses information which impacts on individual privacy. This should in turn reduce the likelihood of the Organisation failing to meet its legal obligations.

10.7 Conducting and publicising a PIA will help our Organisation to build trust with our young people. The actions taken during and after the PIA process can improve our understanding of our young people. There can be financial benefits to conducting a PIA. Identifying a problem early will generally require a simpler and less costly solution. A PIA can also reduce the ongoing costs of a project by minimising the amount of information being collected or used where this is possible, and devising more straightforward processes for staff. Consistent use of PIAs will increase the awareness of privacy and data protection issues within an organisation and ensure that all relevant staff involved in designing projects think about privacy at the early stages of a project.