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"Not everything that can be counted counts and not everything that counts can be counted"

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Useful terms

Output

Outputs are the things you do and activities you undertake to achieve your outcomes. These often contain action words like: *deliver, provide, facilitate, create*.

Outcome

Outcomes are the change or difference you want to make through the work you do. Outcomes should include change words like: *increase, decrease, reduce, improve*.

Impact

The longer-term difference or change that your organisation or project makes.

Outcome indicators

Indicators are the things you measure to tell you whether or not you're achieving your outcomes. Indicators may also be about measuring the steps along the way towards large or long-term outcomes.

Qualitative evidence

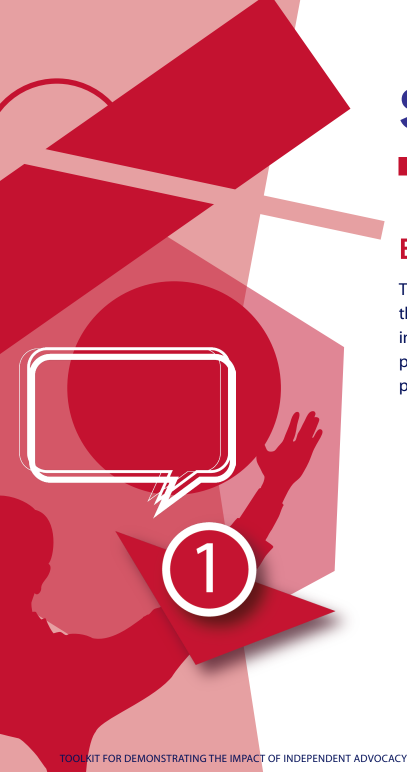
Qualitative evidence is descriptive information about people's experience, perceptions or feelings.

Quantitative evidence

Quantitative evidence is numerical or statistical information: how much? How many?

Baseline information

Evidence you collect at, or before, the start of a project or intervention. Baseline information allows you to measure distance travelled or progress towards outcomes.



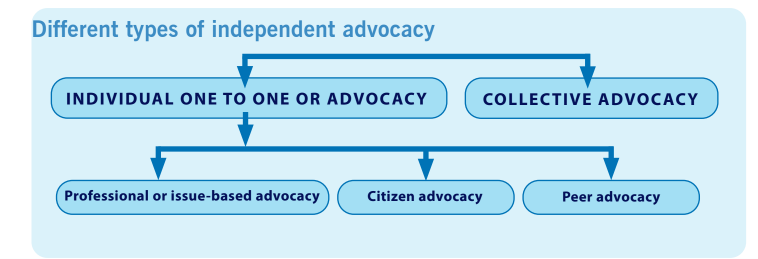
Section 1: Introduction

Background

The Scottish Independent Advocacy Alliance (SIAA) has released this updated version of the toolkit, designed to assist independent advocacy organisations throughout Scotland in demonstrating their impact. The introduction of the national outcomes' framework will provide a means for independent advocacy organisations to reliably report on the impact and positive changes made for individuals and communities in Scotland.

Who is the toolkit for

The toolkit has been developed and designed so that it is relevant to both **individual** and **collective** advocacy.



The toolkit is useful for:

- Board, managers and senior staff working in independent advocacy organisations
 who have a leadership role in designing and delivering work to demonstrate
 impact, as well as cascading the knowledge and relevant processes throughout the
 organisation.
- **Workers** and **volunteers** in independent advocacy organisations who have a role in demonstrating impact by gathering, analysing, and utilising evidence.

Commissioners and funders involved in the provision of independent advocacy. It
will support them in developing evaluation frameworks that ask meaningful questions
in appropriate ways that demonstrate the impact of independent advocacy. This
will, in turn, ensure value for money, robust quality assurance mechanisms and the
ongoing enhancement of independent advocacy.

Defining 'demonstrating impact': outputs v. outcomes

This toolkit provides a practical guide to measuring and reporting the impact of independent advocacy. It offers a step-by-step approach to demonstrate how advocacy partners, groups, other organisations, and the wider community benefit from these efforts. Traditionally, many organisations were only obliged to report on the activities they delivered with their funding, such as the number of people receiving advocacy or hours of advocacy support provided. However, the current emphasis on demonstrating impact requires more than just output reporting. Independent advocacy must also describe and provide evidence of the difference made by these outputs, also known as outcomes.

By collecting clear, structured feedback, independent advocacy organisations can develop in ways that enhance their effectiveness and accessibility. The evaluation process can also identify and promote models of best practice, as well as identify trends, key issues, and gaps in independent advocacy, which can inform policy development and the advancement of independent advocacy more widely.



Demonstrating impact: National outcomes and organisational outcomes

Our toolkit offers support for measuring and reporting on both the Independent Advocacy National Outcomes and individual organisational outcomes. We are collaborating with our members to measure the impact of the Independent Advocacy National Outcomes, using an outcomes framework developed in partnership with our membership.

Independent advocacy national outcomes

One of the struggles, nationally and beyond, in demonstrating the impact of independent advocacy is that independent advocacy organisations do not have consistent outcomes towards which they are all measuring.

The research conducted when developing this toolkit showed that whilst independent advocacy organisations may be very rigorous in collecting evidence of their outcomes and the difference they are making – but that these are not necessarily the *same* outcomes.

This is a collective issue for independent advocacy organisations in Scotland, that makes the SIAA's work at a national level, to promote, support and advocate for independent advocacy and positively influence policy, more challenging. A national dataset will enable SIAA and its members to measure and explain their impact, learn how to improve practices, and inform policy and the development of independent advocacy.

To address this identified gap SIAA underwent a period of consultation with members and agreed three **Independent Advocacy National Outcomes**:



Independent Advocacy National Outcome 1

Independent advocacy enables people to be better informed about their rights, as well as understanding their needs and wishes and the choices they have. It increases their agency and power to choose their own actions freely, as much as possible.

Independent Advocacy National Outcome 2

Independent advocacy supports people to recognise, understand and challenge power imbalances that influence their lives. It enables them to challenge these imbalances, in order to try and realise their rights, needs and wishes.

When someone is unable to take personal action, their advocacy worker will ensure that their rights are upheld.

Independent Advocacy National Outcome 3

Independent advocacy supports people to know that their voices have been heard and their needs and wishes understood.



Organisational outcomes

In addition to the three Independent Advocacy National Outcomes, this pack will also support you to gather evidence and report on any additional organisational outcomes that you feel are important, or that your commissioners/funders require you to evidence.

The toolkit provides information and a range of tools and approaches for gathering, analysing, and reporting on reliable evidence of the wide-ranging impacts of independent advocacy. Doing so will contribute to the development and sustainability of independent advocacy.

"It is important to remember that independent advocacy highlights opportunities, supports people to be more aware of opportunities, choices and their rights, enabling them to make more informed decisions and to become more influential as agents of change. Through broadening horizons and widening understanding of options, independent advocacy enables people to educate themselves and be more active citizens."

Scottish Independent Advocacy Alliance





Section 2: Why evidence the outcomes your organisation contributes to

The benefits of gathering evidence

Independent advocacy organisations are usually required to conduct some form of monitoring and evaluation for their work, but often it is viewed as a necessary activity for the benefit of funders rather than the organisation itself. However, gathering evidence of the difference that independent advocacy organisations make can have a range of benefits for both the organisation and the people accessing independent advocacy.

Internal benefits for the organisation include enhancing practice, informing new independent advocacy development, motivating, and affirming staff, and fostering a climate of critical and constructive thinking. Such benefits contribute to improved independent advocacy for people.

Furthermore, independent advocacy organisations that measure, analyse, and evaluate outcomes can use their evidence to influence local and national policy and independent

advocacy provision. They can communicate more clearly and authoritatively with potential advocacy partners, other professionals, and the media about the value of independent advocacy to individuals and wider society. They can also have a realistic understanding of what the organisation is achieving with people seeking advocacy support and the changes that could be made to strengthen practice. By noticing trends, organisations can address issues proactively.

In addition to these benefits, evaluation enhances accountability to independent advocacy boards, staff, volunteers, and most importantly, the people and groups being supported.

To ensure that organisations are providing the most effective, efficient, accessible, and person-centred, they must critically examine the impact of advocacy interventions on those who use them. This level of accountability is essential to key stakeholders and enables truly transparent organisations. The resources provided in this toolkit are designed to encourage organisations to collect and leverage feedback from advocacy partners or collective advocacy members. By doing so, organisations can make informed decisions on developments to better support those who access independent advocacy.

How to use this toolkit

This toolkit offers a range of flexible tools and approaches – *resources* – which are intended to enable independent advocacy organisations to adopt a realistic and appropriate framework for evaluation in their organisation.

The focus of the tools and approaches within this toolkit is primarily on outcomes – or whether or not the organisation is *making the difference* it aspires to.

There will also be some examples of tools that help organisations learn about how well processes are working (which helps understanding if there are certain groups, or individuals



with particular circumstances, for whom processes need to be adjusted in order to achieve the desired outcome.)

Some of the independent advocacy organisations we spoke with when researching this toolkit are currently asking questions on evaluation forms like 'Did your advocacy partner explain their role?', and 'Was the service easy to use?' These are good examples of questions that will help us understand how well our processes are delivering for us – but they don't tell us about the differences that advocacy is making – or outcomes.

Whether your organisation is new to gathering evidence of the differences that its make, or seeking to refresh approaches it is already using, our recommendation is that having a read through the pack in its entirety will ensure that you get the most from it. We will be clarifying important terms, providing useful examples and top tips – as well as providing tools and approaches, in the Resources section, that you might use.

While it is most often managers or senior staff who have responsibility for organisational 'monitoring and evaluation', there is much to be gained from promoting these strands of work as a shared responsibility amongst all staff and volunteers. Creating a culture in which enquiry, critical thinking and using evidence to inform practice and development will strengthen any organisation; as well as result in excellent provision those accessing independent advocacy.

While this toolkit offers guidance, tools, and approaches, it is important that each organisation discusses and clarifies the parameters of their approach to monitoring and evaluation, as there is not one size that fits all. While the principles and practices will be similar, the scope, depth and ways that evidence is used are likely to vary.





Section 3: What are outcomes

Organisations are accustomed to reporting on activity in terms of outputs. However, in recent years, there has been a growing demand for outcome reports showing the impact that independent advocacy makes for people and groups. This has become even more important given the Scottish Government's shift towards an outcome-focused approach.

Outcomes are the changes your organisation makes to the lives of the people it works with. Outcome statements often include words that indicate a change or difference e.g. increase, reduce, expand, enable, improve, sustain.

Outcomes may be things like knowing that one's wishes have been understood, having a decision made by an agency successfully overturned, or feeling safer in one's home.

When evaluating outcomes, it's essential to differentiate between short and long-term goals. For instance, if an organisation decides to alter its offering based on collective advocacy, this might be viewed as a short-term outcome. However, if the actual service delivery changes, it could be considered a longer-term outcome to continue striving towards. On an individual level, comprehending one's rights is a short-term outcome, while working to guarantee those rights are respected is a longer-term objective.

Outputs are the things you do and activities you undertake to achieve your outcomes. They often start with words that indicate activity e.g. deliver, offer, provide, facilitate, create.

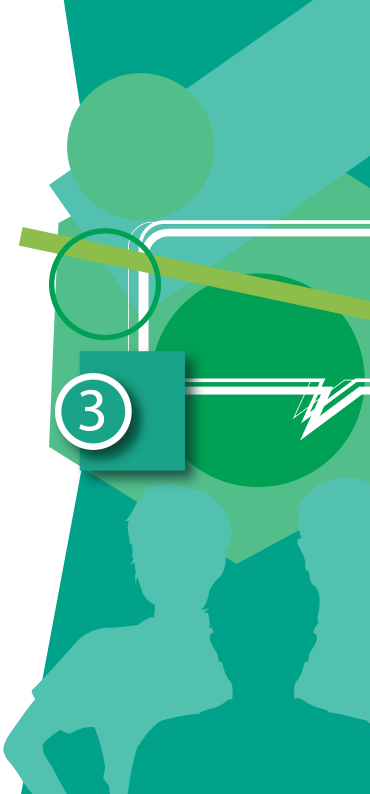
When organisations describe the services they offer – providing independent advocacy support, giving information, attend meetings – they are describing *outputs*.

Outputs are what we do, and outcomes are the differences that occur because of what we do.

While the SIAA expects that its members collect evidence in relation to the three Independent Advocacy National Outcomes highlighted in the introduction, so that it can use Scotland-wide evidence to strengthen its work on a national basis, it is likely that individual independent advocacy organisations will have additional outcomes that they need, or wish, to evidence.

What matters is that independent advocacy organisations are clear about *the differences* that will be made because of their outputs. Once they are clear about these differences, they can create strong outcome statements (the three Independent Advocacy National Outcomes are examples of this).

Once the outcomes statements have been agreed within the independent advocacy organisation, thought can be given to how to go about gathering evidence to demonstrate that they are being achieved.





Section 4: How will we know we are making a difference Understanding outcomes and outcome indicators

Once an independent advocacy organisation is clear about the difference it is seeking to make (*outcomes*), and the activities it will deliver for the purpose of making such a difference (*outputs*), it is time to think about the evidence that will demonstrate success – or **outcome indicators.**

Each outcome will probably have a range of potential 'indicators'.

For instance, many independent advocacy organisations support people to participate more in situations that affect their lives (outcome) – through providing individual and collective advocacy (outputs).

In addition to answering 'yes' on an evaluation form that asks whether people believe

that advocacy resulted in them participating more in situations that affect their lives, an advocate might hear someone describe:

- An improved relationship with their health/care provider
- Feeling more confident, generally, to raise concerns when they have them
- Using more assertive behaviour with family or friends

All these statements are legitimate 'evidence' that someone is participating more in situations that affect their lives. When we spend time deepening our understanding of the outcomes we are working to achieve with people, it is much easier to gather evidence, conditions, and behaviours because we are noticing it. This will be a useful point for managers/senior staff to emphasise when supporting staff to notice and collect evidence. During our research with SIAA members whilst developing this toolkit, we heard that becoming aware of the 'informal' feedback that is being offered can become a real motivator for staff and volunteers.

When establishing outcome indicators, it can be useful to ask questions such as:

- What could success look like?
- What types of things might be happening for a person if this was true?
- How have other people described this to us?
- What kind of changes would we expect a person to describe to us that would suggest they have... (achieved stated outcome)

While people using independent advocacy organisations generally come with a specific purpose in mind around which they are seeking advocacy support (and which generally aligns to one of the national outcomes), they may have some additional personal outcomes such as developing their confidence to speak in official meetings, being clear about what



they want – and asking for it, or managing their emotions more effectively in official settings.

Some of the tools in the Resources section make it easy for advocacy partners/ collective advocacy members to identify their own outcomes to work toward, in addition to those that the organisation is seeking to evidence. How, when, and why you might do this will be explored further in the Resources section of the toolkit.

"When we write our outcomes, we should include the "who" in the outcome. For example, "care experienced young people have more positive life chances". That way we know who we intend to make a difference with and for, and who we should ask for feedback from when measuring the outcome.

Indeed, the people the outcomes are for should really be involved in writing them in the first place. And people should be at the heart of measuring the outcomes approach.

They – not (just) the stats – tell us whether activities are making a difference and why.

That in turn means that the outcomes approach requires a rich tapestry of evidence – research, practitioner experience, lived-experience evidence; a mix of stats and stories".

Steven Marwick, CEO, Evaluation Support Scotland (2019)





Section 5: What counts as evidence

Qualitative and quantitative evidence

There are two main types of evidence – qualitative evidence is descriptive (e.g., case studies) and quantitative evidence is numerical (e.g., statistics). These are explained in more detail below.

For independent advocacy organisations, the most engaging evidence to collect as evidence that outcomes are being achieved will be qualitative – most often relating to changes in thinking, behaviours, and feelings. Qualitative evidence is what we hear when people tell their stories – beyond 'yes and no' descriptions of whether intended outcomes have been achieved. Consider, for a moment, all the ways in which people's lives have changed because of their wishes being heard and understood. Each of those stories is providing qualitative evidence that this important outcome has been achieved. That type of evidence is rich and varied.

However, *quantitative* evidence is also useful when demonstrating that we are achieving outcomes. Questionnaires that ask whether someone believes an outcome has been

achieved can be used to produce quantitative evidence. When we report this type of evidence, we generally do so numerically. For example:

'85% of people who received advocacy report that they were supported to put their point of views across'. Or '95% of people who received advocacy report that they have gained in understanding of their rights.'

Evidence which can be collected in relation to outputs is also likely to be quantitative – relating to numbers – a number of advocacy hours, the number of advocacy partners in particular age or post-code categories, the number of information sessions delivered, etc.

Quantitative and qualitative evidence have the most impact when they are used together. For example:

"In 2019, we delivered 2,465 hours of advocacy support to 237 individuals, 93% of whom reported that their voices were heard, and their wishes understood. They told us that:

'Having the support is invaluable to me because I don't think I could have gone to my DWP assessment on my own.'

'My advocate expressed my views at tribunal on my behalf'.

'With support I was able to attend meetings regarding my daughter. I was confident to put my views across and I now have contact again.'

'I was able to have my views on where I live communicated through my advocate'.

Organiations often utilize a combination of quantitative and qualitative data through case studies, which can be applied in various contexts. These case studies are beneficial for funders, individuals considering advocacy support, annual reports, and general promotion of the significance of independent advocacy.



Determining which evidence to collect, as well as the approach and timing for collecting it, are crucial choices to make. By creating an organisational framework, all staff can work together in a consistent and efficient manner when engaging with individuals who have received independent advocacy to provide feedback. This helps ensure that the organisation is accomplishing its intended outcomes and allows for the gathering of valuable information on processes.

What counts as evidence?

When we think of evaluation, we tend to rely on direct feedback mechanisms such as questionnaires and interviews to provide evidence. However, while these are effective tools, they can be resource-intensive and require significant planning. It's vital for independent advocacy workers to think of evaluation as something integrated into their day-to-day practices, rather than a separate task. This way, they can consider all the forms of evidence that already exist – including notes, thank-you messages, and positive changes in appearance, body language and behaviour – that might go unnoticed and unrecorded.

Other examples of valuable sources of evidence include stories from people using independent advocacy and observations from friends, relatives, and other professionals. While direct feedback from advocacy partners is the most important source of evidence, observations from others are also valid. By being clear about the desired outcomes and the indicators required to demonstrate success, workers can more easily recognise the relevant pieces of evidence.

In this toolkit, we highlight the importance of integrating evidence collection into existing systems and practices, rather than adding extra work for workers and volunteers. Once the desired outcomes and evidence types are established, workers can decide on the best methods for collecting, recording, and storing the evidence.



While getting direct feedback from someone who has accessed independent advocacy is the most important source of evidence, observations from others are also valid sources of information. It might not be possible to get feedback from the advocacy partner, when providing non instructed advocacy and therefore it is important to gather feedback from others involved in the situation, such as unpaid carers.

What we see and hear, as well as what others (family, friends, other professionals) tell us about changes they have noticed is all data. All, useful evidence.

Throughout this toolkit, you will notice a theme – that recording evidence should, as much as possible, be integrated into existing systems and practices, rather than something additional that workers and volunteers need to make significant extra time for.

Once you are clear about the outcomes you want to be able demonstrate and have thought about the types of evidence that you might see or hear that would show/tell you they are being achieved – you can decide how to collect, record and store it.





Section 6: Collecting evidence: What, when, how and where

The Toolkit's guidelines for collecting and reporting evidence are designed to be adaptable. This means independent advocacy organisations can tailor the process to fit their specific needs. Some organisations may choose to focus solely on the necessary requirements while others may prefer to gain a more comprehensive understanding of the process. Ultimately, this approach accommodates a wide range of preferences and priorities.

What evidence to collect

There are two kinds of evidence which can be useful to collect – and some which is often required by funders and commissioners.

- that which relates to the difference the organisation is making
- that which relates to how it goes about making that difference

The first is evidence which relates to **outcomes** – the difference the organisation is making

to the people receiving its services. This is the type of evidence that funders often require.

Through gathering, and thinking about, this type of evidence, you will get clearer about outcome indicators; what you see and hear that tells you an outcome has been achieved.

The second, is evidence to show that the organisation is functioning in the most effective and efficient way possible. This kind of evidence relates to **processes** – how the organisation delivers what it sets out to achieve.

In practice it is useful to gather both outcome and process data at the same time, where possible – both to save time and to avoid over consulting people.

Most of the independent advocacy organisations we spoke with are already gathering this type of evidence, in relation to service quality and how it is delivered. For example, they are asking questions such as:

- Did your advocacy worker listen to the things you had to say?
- Did your advocacy worker explain their role, and how they could help?
- Did you find our service was easy to use?
- Are you clear about how to make a complaint, if you are unhappy with our service?

These types of questions do not tell us whether we have delivered the outcome we are working toward – but they can potentially give us a lot of useful information about how to deliver our best for people who are seeking advocacy support.

Questions about processes are useful and particularly important if something about the organisation changes – such as opening times, or the advocacy provision that are available – so that we can learn about the affect this change is having.



This type of question might also be asked only for a period, in specific circumstances, or of certain people (specific segments of the client group who would be most affected by a change, etc.) when we are trying to better understand something about our organisation.

For example, if we are trying to improve the experience of being on a waiting list, we might ask only people who are waiting about their ideas for improving this experience.

The best way to decide on the type of evidence or data you need to gather is to refer to your **outcomes and outcome indicators**. These will give you clear guidance as to the kind of information you need to collect. The Resources section of this pack will be useful for determining how often you collect it, and by what means.

Qualitative or quantitative data?

Remember that you want to collect both *qualitative* (usually relating to thinking and feeling) and *quantitative* (things we can count) evidence.

Stories and quotes are *qualitative* evidence.

Self-assessment 'rating scales' like Outcome Stars are very useful for providing quantitative evidence e.g. 'When asked to rate 'do you feel that the services you are engaged with understand your wishes?' at their intake appointment, 15 people who scored themselves 4 (or less) out of 10 were scoring 7/8 out of 10, at the end of their advocacy support.'

Closed questions also provide good quantitative evidence. Closed questions are those which require a 'yes' or 'no' answer. Open questions often start with *how, who or why* and



require a more detailed answer. These are the type of questions where a text box, or space to write, is offered.

For example:

'Do you think you benefited from advocacy?' Yes/No –provides evidence that enables the organisation to make statements like '97% of people asked reported that they have benefitted from advocacy.'

These can be paired with an 'open' question providing qualitative evidence

'If you answered 'yes', what difference do you think independent advocacy has made to you?'

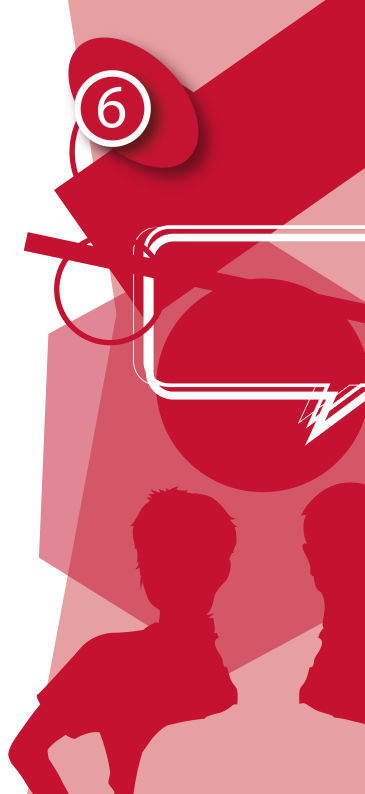
Make it Easy: Thinking about the evidence that you need to collect, craft a small set of questions that you believe will provide it (remember to only collect what you need!) Make these a blend of open and closed questions, ideally that can be included in the same form, on-line questionnaire, phone call or focus group interview (all methods that are detailed in the Resources section).

When to collect evidence

There is no 'right' time to collect evidence, but there are useful things to keep in mind when you are considering when might be the best time(s) to do so.

In order to assess whether a change has happened – in other words, whether independent advocacy support has made a difference – it is advisable to start by collecting information from the advocacy partner/group member at an early stage.

This is because, in order to measure the distance travelled accurately, you need to know where the journey began. This information is often referred to as 'baseline information', or simply a 'baseline'.



You don't need to start collecting evidence the first time you meet the advocacy partner – in fact, the majority of independent advocacy providers we consulted with commented that trying to gather information in a first meeting frequently feels like the wrong thing to do. Most of them, however, reported that they start to gather baseline evidence within the first three visits, once the relationship feels established enough to ask someone to engage in this way.

Having an organisational policy that this type of information will be gathered **within the first three meetings** provides enough flexibility for independent advocates to decide when it is best to start the process based on their judgements within each individual independent advocacy relationship. In some cases, this might happen naturally within the first meeting, whilst for others it might make sense to wait until the relationship has been established.

The baseline information collected needs to relate to the improvements your independent advocacy organisation is trying to make, which is likely to go beyond the types of questions that you ask a person at the point of offering advocacy. However, it is possible to include some of the key questions as part of an initial interview if you judge it appropriate to do so.

A number of the tools contained in the Resources section are useful for collecting baseline information. There are several examples of rating scales that make this very easy and are all useful tools for collecting information at the beginning, and then again at stages throughout a person's independent advocacy journey so that changes can be measured.

Asking some questions at the end of an advocacy partnership or life of a collective advocacy group is essential – and this is what we found that most organisations are



doing consistently. The weakness of this approach is that once the situation has been resolved the advocacy partner doesn't want to think about the difficulties they previously faced and has moved on with their lives. Also sending out printed forms for advocacy partners to complete may present problems for those with literacy issues and might not be an accessible format for everyone. Some advocacy partners might need the help of an independent advocate to complete the form itself. Unsurprisingly, most organisations reported poor returns when they send out printed forms – generally less than 20%.

Therefore, it is important to consider the method of collecting feedback and having a minimum return rate in mind. The other disadvantage to asking for feedback only at the end of the relationship or intervention is that if the advocacy partner's desired outcome is not achieved that might cloud the way they feel about the way the advocacy was delivered. It is perfectly possible for an effective independent advocate to follow all the principles of independent advocacy and the code of best practice and still not achieve the desired outcome of the group or individual being supported. It might be difficult to separate the disappointment with the outcome from the process, which may have been 'by the book'.

Comparing baseline evidence with that which is reported at the end allows us to measure progress and to report confidently on the difference that independent advocacy makes. Again, tools like the various rating scales, which include some that are picture based, in the Resources section make it easy to compare 'before and after advocacy'.

Many of the independent advocacy organisations that we spoke with during our research are only seeking to gather data at the start and the end of the advocacy process, as a way of evidencing that key outcomes have been achieved. *This is absolutely good enough!*



How to collect evidence

In the Resources section, there are useful tools and other ideas for collecting evidence, including a number of 'rating scales'. Because scales – which come in a variety of forms – make measuring progress so straightforward, they are worth a specific mention in this section of the toolkit.

A 'rating scale' is used to respond to a statement by selecting a point, often on a numerical scale, that one believes best represents their position in relation to the statement. For example:

'I am confident that my wishes have been understood'.

1 (not at all) 2 (unsure) 3 (somewhat confident) 4 (fully confident).

As well as being uncomplicated to use, rating scales are also useful as a consistent means of gathering the same evidence from an individual or group over time. There are examples of different rating scales of the Resources section of this toolkit.

Rating scales are useful because:

- They provide a clear 'baseline' to measure against at any future stage
- They can relate directly to organisational outcomes
- They can be used with some flexibility as each advocacy relationship is unique
- They can be used to establish an individual's own goals/aspirations (in addition to those the service is seeking to measure)
- They have potential to strengthen people's personal resilience, as they reveal 'real data' that reflects progress across time



• Funders often like them because they can provide a 'snapshot' of a service – charts and graphs representing grouped feedback, along with quotes, are ideal for sharing both quantitative and qualitative evidence.

It is important to explain to staff and volunteers to be confident about *why* information is being collected, and *how* the information will be used. When they are clear, they are able to confidently explain this to those who are being asked to share their information.

It should also be made very clear to advocacy partners and collective advocacy members that this information is in NO WAY intended as a means of comparing one person's progress with another's.

As mentioned earlier, it is important that every person understands that while the independent advocacy organisation needs and values this information, the advocacy partner/group member is not obliged to provide it – it is done so voluntarily, for the purpose of developing and sustaining the organisation.

To maximise the flexibility of this toolkit, and in acknowledgement of the fact there can be no 'one size fits all' approach, a range of tools and approaches for gathering evidence are explored in the Resources section. It is advisable to spend some time talking them through as a team, exploring questions like:

- What tools look like they would work well with our advocacy partners or groups?
- Are there any that we think are not the tools for us to use? Why?

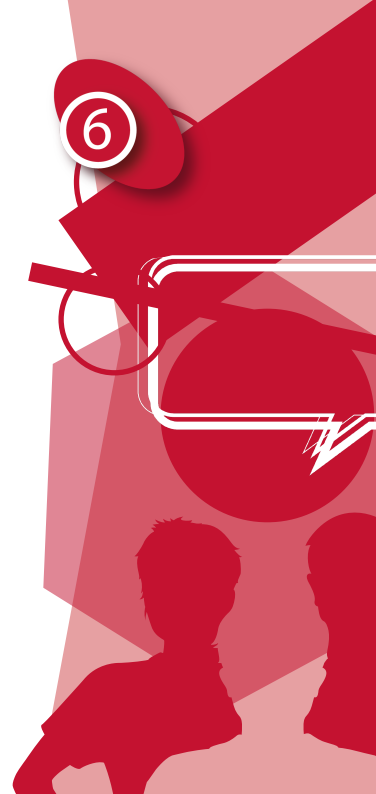


It will also be important to discuss the methods which may be most appropriate for different types of clients, at different stages of a partnership or collective group, and in particular settings.

For instance – what might be the best method for people with literacy issues or with specific communication needs? Children and young people? Those who use English as a second or third language? The best methods in 1:1 versus group settings?

To develop a successful evidence-based approach for your independent advocacy organisation, it is essential that staff work collaboratively to customise the approach over time. The gathering and use of evidence will be most effective if diverse perspectives are considered in the process of drawing conclusions from the data. This includes input from staff, volunteers, advocacy partners, and collective members. Everyone brings a unique viewpoint on what the evidence is conveying.

To ensure consistency of approach across the organisation, staff must feel confident in their ability to select appropriate tools and introduce the evaluation tool. It is recommended that all staff receive adequate training to generate this confidence. Continued discussion and clarification amongst staff will help maintain consistency and ensure ongoing success. Additional resources on this topic can be found in the Resources section.



Most of the independent advocacy organisations we visited told us that gathering evidence in a systematic way feels clumsy, and even uncomfortable, to begin with... but, critically, they said it gets much easier with practice. It may be a big learning curve for staff – or not. It is useful to think together about how to weave it into the 'usual' way of working as much as possible; and avoid making it yet another thing to do!

The key is to gather only the information you are going to use, and to record evidence as you go along.

It is useful to ask advocacy partners and collective members what questions they think the organisation should be asking, and when they want to be asked. They bring an important perspective on what your organisation needs to better understand, in order to achieve its potential for all involved.

There is no 'best way' to collect evidence. Because independent advocacy services are different, just like each person, it is for each organisation to decide when and how to gather evidence. This toolkit should provide everything a service needs to get started, or to refine and develop its current approach.



Where to collect evidence from

As previously mentioned, evidence is constantly present, however, it may go unnoticed and unrecognised. Once you have established your desired outcomes and indicators, you will be better equipped to identify the many ways in which evidence is readily available to you. Evidence can be obtained from a variety of sources, including advocacy partners, collective advocacy members, and their respective support systems, whether that be family or friends. Additionally, evidence can be gathered from both paid and unpaid staff, as well as other professionals. These professionals may include referrers, partner organisations, or anyone granted permission to provide feedback. It is important to note that much of the evidence you seek may already be available, often unknowingly provided by those around you. The key is to be observant and take note. Resources such as paper or electronic forms, phone interviews or focus groups can also be utilised to obtain necessary evidence.

In the case of thank you notes/emails, When you hear someone comment about a difference in themselves or you observe changes in their behaviour or appearance, that you believe may indicate that an 'outcome' has been achieved its always important to have a conversation with the person to check and confirm what you have noticed.

That might sound something like this -

Joe, I hear you describing that you are being more assertive with your CPN, asking for what you need. It sounds like she is really hearing what you are wanting to happen. Does it seem like that to you?



Evidence of outcome: People know that their voice has been heard, and their wishes understood.

Additionally, we might have agreed indicators that can be observed – such as a change in behaviour or appearance. If we notice a change in our advocacy partner or member, it is a good excuse to start a conversation about it – do they have any sense that it is related to advocacy?

If the answer confirms that what has been heard or observed reflects an outcome being achieved, then the next thing to do is to ask for permission to record it – as evidence.

Remember that you may need to explain why the organisation needs evidence, and how i will likely be used.

Storing evaluation/outcome information

When it comes to advocacy organisations, striking a balance between respect for individual wishes, maintaining confidentiality, and demonstrating impact is crucial. Administrative systems must be GDPR-compliant, and data must be considered within this framework. If you require assistance in this area, refer to helpful websites such as https://gdpr.eu/ and Data protection: The Data Protection Act - GOV.UK (www.gov.uk).

This toolkit is designed to help independent advocacy organisations in Scotland confidently gather evidence of their impact on the three Independent Advocacy National Outcomes, as well as other important factors. A list of additional useful links and resources is included at the conclusion of the toolkit, should your organisation seek to further refine its evidence-gathering in impact measurement.





Resources

This section of the toolkit offers a range of tools for gathering evidence, and some thoughts on ways in which this evidence can be used to strengthen independent advocacy organisations.

Everyone who is asked to complete a questionnaire, or other tool, should understand why they are being asked for their opinions (why the organisation is collecting the information), and what handing over their information will mean (how it will be used).

They should also be clear that – while the organisation values the information – they are under no obligation to share it.

The outcomes framework

SIAA and its members have collaborated to create a practical outcome framework that breaks down the three national outcomes to measure the impact of independent advocacy in Scotland.

Members from collective advocacy groups, individual advocacy and citizen advocacy came together to make certain the outcomes framework would be suitable for every model of independent advocacy and indicators relevant for all groups and people. The framework features clear indicators and **suggested** evaluation methods, which are only examples of how data could be gathered rather than a rigid list. Independent advocacy groups may use various techniques and tools to collect data. The infographic below shows how the principles and standard which underpin good practice impacts on:

- Advocacy partners and collective advocacy groups
- Other organisations and services
- The wider community

Principles, standards and outcomes

R

Principles

Standards

1. Independent advocacy stands up to injustice, discrimination, and disempowerment.

Independent advocacy recognises power imbalances or barriers people face and takes steps to address these.

Independent advocacy enables people to have more agency, greater control, and influence.

Independent advocacy challenges discrimination and promotes equality and human rights.

2. Independent advocacy is loyal to the people it supports and stands by their views and wishes

Independent advocacy follows the agenda of the people supported regardless of the views, interests, and agendas of others.

Independent advocacy must be able to evidence and demonstrate its structural, financial, and psychological independence from others.

Independent advocacy provides no other services, has no other interests, ties or links other than the delivery, promotion, support and defence of independent advocacy.

3. Independent advocacy ensures people's voices are listened to and their views are taken into account.

Independent advocacy reduces the barriers people face in having their voice heard because of communication, or capacity, or the political, social, economic, and personal interests of others.

Independent advocacy recognises and safeguards everyone's right to be heard.

These standards impact on



Advocacy Partners/Groups outcomes

People are better informed of their rights.

People are more able to express their views

People have their views and wishes taken into account.

People have more choice and control in situations which affect their lives.

Other organisations and service outcomes

Improved services with greater equality of access.

People are treated with more dignity and respect.

Independent advocacy credibility and legitimacy is recognised and fully understood.

Independent advocacy helps to defend and uphold human rights.

Wider community outcomes

People in Scotland understand what independent advocacy is and can access it if required

Reduction of stigma in communities.

Empowered community with strong connections.

Human rights upheld for all, strong community voice.

Independent Advocacy National Outcome Statement 1

Independent advocacy enables people to be better informed about their rights, as well as **understanding their needs and wishes and the choices they have.** It increases their **agency and power** to choose their own actions freely, as much as possible.

Independent Advocacy National Outcome Statement 2

Independent advocacy supports people to **recognise, understand and challenge power imbalances** that influence their lives. It enables them to **challenge these imbalances**, in order to try and realise their rights, needs and wishes.

When someone is unable to take personal action, their advocate will ensure that their rights are upheld.

Independent Advocacy National Outcome Statement 3

Independent advocacy supports people to know that their voices have been heard and their needs and wishes understood.

Advocacy partners or groups

Advocacy partitions of groups					
Outcome	Indicator	Evaluation			
People are better informed of their rights.	 People take part in meetings/decision making processes. People are having conversations about their options. People are talking about their rights. People are given accessible information about their rights. People are given a range of ways to make choices. 	 Questionnaires Self-rating scales 1:1 interview Focus groups Informal feedback Observation Case notes Statistical information 			
People are more able to express their views.	 People tell us they are more supported. People have increased confidence in expressing their views. Less need for independent advocacy. People supporting each other. People are making choices. People contribute at hearings/meetings. Greater number of people are participating in important meetings about them. Increase in evidence of self-advocacy (or supported to self-advocate). 	 Questionnaires Self-rating scales 1:1 interview Focus groups Informal feedback Observation Case notes Evidence of different communication methods 			



Advocacy partners or groups

Outcome

3.

People have their views and wishes taken into account. Indicator

- People are given a range of ways to make choices.
- People tell us they are more supported.
- People are given accessible information about their rights.
- People are more involved.
- People can describe what they want to happen.
- Decisions are taken which reflect the full rights of a person who is unable to act on their own behalf. (Non instructed advocacy)
- People contribute at hearings/meetings.
- Others understood what the person/group wishes to happen.
- Active consideration is given to the true wishes of a person/group by the professional(s)/others in a meeting

Evaluation

- Questionnaires
- Self-rating scales
- 1:1 interview
- Focus groups
- Informal feedback
- Observation
- Case notes



Advocacy partners or groups

•

Outcome

4.

People have more choice and control in situations which affect their lives

Indicator

- People have increased participation in decision making processes.
- People have their rights safeguarded (non instructed advocacy)
- Decisions are taken which reflect the full rights of a person who is unable to act on their own behalf
- People are given a range of ways to make choices
- · People are more involved
- Increase in evidence of self-advocacy (or supported to self-advocate)
- People are making choices
- People contribute at hearings/meetings
- People felt more powerful in conversations about them and their lives
- People have a greater sense of self identity and autonomy
- People are more empowered

Evaluation

- Questionnaires
- Self-rating scales
- 1:1 interview
- Focus groups
- Informal feedback
- Observation
- Case notes

	Other organisations or services			
Outcome	Indicator	Evaluation		
People are treated with more dignity and respect	 People are given a range of ways to make choices People express they feel more supported People are given accessible information about their rights and the options available to them People are more involved People contribute at hearings/meetings People express a greater sense of self identity and autonomy 	 Questionnaires Self-rating scales 1:1 interview Focus groups Informal feedback Observation Case notes 		
Independent advocacy credibility and legitimacy is recognised and fully understood	 Independent advocacy is connected strategically and locally to Boards, forums, steering/working groups as an equal partner whilst maintaining their independence Evidence of systems change/policy in response to input from independent advocacy organisations/groups Increased referrals Increased funding for all models of independent advocacy 	 Strategic Plans Local Advocacy Plans Local Policy Evaluations Advocacy Map Local reporting National outcomes report Annual reports 		

Other organisations or services

2 11 2 1 8 1 1 1 2 1 1 1 2 1 1 1 2 1 1 1 2 1 1 1 1 2 1 1 1 2 1					
Outcome	Indicator	Evaluation			
3. Independent advocacy helps to defend and uphold human rights	 Advocacy organisations/groups gather and use data on the number of people in communities facing the same issues or difficulties Advocacy organisations/groups shine a light on local and systemic issues Advocacy organisations/groups contribute to using a rights-based approach in local communities Advocacy organisations/groups have raised awareness of human rights Local communities have an increased awareness of stigma and discrimination Local communities have increased empathy and understanding of everyone's rights 	 Evaluations Advocacy Map Local reporting National outcomes report Annual reports Strategic Plans Consultation responses Training in Human Rights 			
4. Improved services with greater equality of access	 More people can access other services at the right time Increased awareness raising sessions in community Other services are clear about the role of independent advocacy and make referrals Less people presenting to crisis services or in crisis 	 Questionnaires Self-rating scales 1:1 interview Focus groups Informal feedback Observation Case notes 			

Wider community

Outcome

1.

People in Scotland understand what independent advocacy is and can access it if required **Indicator**

- People and communities know independent advocates have the values, skills and knowledge required for the role
- More people can access services at the right time
- · Increased awareness raising sessions in community
- Receiving more appropriate referrals
- Less people presenting to crisis services or in crisis

Evaluation

- Questionnaires
- Self-rating scales
- 1:1 interview
- Focus groups
- Informal feedback
- Observation
- Case notes
- Evaluations
- Advocacy Map
- Local reporting
- National outcomes report
- Annual reports
- Websites
- Publications
- Local Advocacy Plans

Wider community

Outcome Indicator Evaluation Questionnaires • Wider communities grow in awareness of stigma and discrimination Self-rating scales • Multiple advocacy organisations gather and use data on the number Reduction • 1:1 interview of people in communities facing the same issues or difficulties of stigma in Focus groups • Independent advocacy organisations/groups shine a light on local and communities Informal feedback national systemic issues Observation Case notes Evaluations Advocacy Map Local reporting • National outcomes report Annual reports Advocacy Plans



Wider community

Outcome Indicator Evaluation

3.

Empowered community with strong connections

- Strong collective advocacy voice within communities
- Local services and policy influenced by independent advocacy
- Communities have increased knowledge of their rights and how to access them
- People are more connected to their local communities through citizen advocacy
- Local communities value the contribution of people who are seldom heard
- Access to justice

- Advocacy map
- National outcomes report
- Inclusion of independent advocacy in policy and legislation

4.

Human rights upheld for all, strong community voice

- Wider community have empathy and understanding of everyone's rights
- · People have their rights safeguarded
- Wider communities are aware of their right to independent advocacy and how to access it
- People are supported to decide whether to use independent advocacy or not
- Strong collective advocacy voice within communities
- Wider policy and legislation are influenced by independent advocacy
- Access to justice

- Advocacy map
- National outcomes report
- Inclusion of independent advocacy in policy and legislation
- Local Advocacy Plans



Once you know what you are trying to measure (outcomes), you are able to select the best tool to use for gathering evidence.

The tools and approaches included here have been selected for their flexibility – they can be used to gather a vast range of information/evidence, from a range of stakeholder groups.

As mentioned in the previous section of the toolkit, there are a variety of ways of gathering evidence. Much of it can be done as part of day-to-day work practice, and some of it will require additional effort.

Methods for gathering evidence are wide-ranging, and should be selected in relation to the capacity of your organisation to undertake them, as well as the needs and preferences of the people you are working with. The simpler it can be made, the more likely it is to happen!

Once I have a tool, who do I use it with?

As mentioned in the previous section of the toolkit, there are other useful 'sources of evidence' in addition to the advocacy partner/collective group member. Friends and family, other professionals, the advocacy worker or volunteer, and other collective group members are all good sources to consider.

The tools in this section are flexible, and can be used with most people. In addition to the table at the beginning of this section related to choosing tools, each example in the toolkit clearly indicates who it is recommended for use with – advocacy partners, family members, advocacy staff/volunteers, and other professionals.

When you know what you what you are trying to measure (outcomes), you can then think about who the potential sources of evidence are – in addition to the advocacy partner or collective group member. You will need to be sure to ask the right questions of the right people – so creating a questionnaire, or other tool, with each different audience in mind is essential. What can you learn from family members? What might you learn from other professionals who signpost people to your organisation – or those who experience the impact of advocacy, through their experience of your work?

While there is some information you will want to gather consistently from certain groups,



such as advocacy partners/collective members, there may be other information that you choose to gather from a 'sample' of people who have received advocacy – or are otherwise connected to the process, such as other professionals.

For instance, a 'sample' of people might be selected for more in-depth analysis, for 1:1 interviews, or to act as a focus group.

Sampling is often done in one of three ways:

- Randomly for instance, choosing every 10th person on the database
- Deliberately choosing individuals who represent a range of characteristics e.g. issue with which they require advocacy, geographical location, age, length of time a service was provided, etc.
- Voluntarily where you issue an open invitation for anyone who is interested to join a focus group, have a telephone interview, or complete a questionnaire.

Once again, it will be for the independent advocacy organisation to decide how much, or how little, it has the capacity to do. We hope that, for all of you, there will be tools in the following section that can strengthen your efforts.



Tools for collecting evidence of independent advocacy's impact Questionnaires

Recommended for use with: Anyone

We heard that questionnaires are the most common tool that independent advocacy organisations use for collecting evidence; and that the return rate for paper-based questionnaires, and therefore the amount of *evidence actually gathered*, is – generally speaking – quite low.

We heard some providers are noticing they get a better response when they send an electronic survey link, or have the form available on the advocate's phone, or a tablet; so that it can be completed in person, rather than posted or emailed.

We also heard that some independent advocacy organisations have found that asking a partner to fill in a final questionnaire in the meeting BEFORE the final one appears to generate richer information than filling it in at the very last meeting. Reflections on the experience of working with an advocate/collective advocacy group are likely to be reported in greater detail when considered separately from any 'results' that have been achieved.

Questionnaires are a means of review, and are particularly useful when seeking to assess an independent advocacy organisation's processes. They can also be useful for evidencing outcomes, but can only do so when they use a combination of *open and closed questions* (see example questionnaires below.)

Asking for simple details like age range and postcode offers additional possibilities for generating quantitative data (things you can count)— as you can identify and report on findings for specific age categories, or people living in certain neighbourhoods.

For example "80% of people between the ages 41-55 rep ort that they are more confident to express their views in a range settings, as a result of independent advocacy – but that is reported to be true for only 67% of people between the ages of 16-25."

While questionnaires are a useful and generally easy way to gather evidence, it is worth putting time into crafting the right questions for your independent advocacy organisation's needs. In addition to the outcomes you are trying to achieve, there may be specific things you want to understand better.

In this section, there are lists of useful questions to consider, for different stakeholder groups. Your questionnaire should contain the questions that you need answered, in order to give you the evidence you require – and any others that will support your organisation to learn about how it can deliver its best.

The first example questionnaire is based on one shared by Angus Independent Advocacy, intended for advocacy partners. Notice the blend of open and closed questions – enabling both quantitative and qualitative data to be gathered, and reported on. You will also notice



that some questions relate to outcomes, and others to processes.

On this questionnaire, questions like 'what worked well', and 'what could have been better' are asking about processes. The question about whether or not someone believes that independent advocacy enabled their voice to be heard more clearly, is a question about outcomes.

Questionnaire: Advocacy Partner

How would you rate the support you received from Angus Independent Advocacy, on a scale of 1(low) to 5 (high)?

1 2 3 4 5

What worked well for you?

3 What could have worked better?

4 Please complete the following sentence:

"Independent advocacy enabled me to ..."

What do you believe changed, because of your advocacy support?

Do you believe that independent advocacy support enabled your views and wishes



	learly understood, than if you hadn't had independent advocacy Yes/No (please circle)				
Why?					
Would you re	ecommend Angus Independent Advocacy to a friend?				
Yes/No (plea	se circle)				
Why?					
•	an you offer any other feedback, that might help us to improve?				
Age range: 1	8-25 26-40 41-55 56-70 70+				
Postcode:					

The following questionnaire comes from Partners in Advocacy (Glasgow), designed for use with children. Respondents are given a sheet of red/amber/green face stickers, like the pictures at the top of the page, to place in the boxes as means of giving feedback on their experience of independent advocacy.



We have chosen to include the following questionnaire, also from Partners in Advocacy (Glasgow) – as it shows the potential to use one questionnaire to learn about a partner's experience both *before* and *after* experiencing independent advocacy.

HOW WAS ADVOCACY FOR ME?



	QUESTION	YES	NO	DON'T	Tell us more if
				KNOW	you like
	Did your				
	Advocacy				
	Worker				
	explain how				
	they could				
	help?				
` @	Did your				
@	Advocacy				
, ,	Worker				
	listen to				
	you?				
	Did				
──¥ YOUR	Advocacy				
YOUR VOICE MATTERS.	help you to				
MAITERS.	have your				
	voice				
	heard?				
	Was there				
3	anything				
	you didn't				
U	like about				
	advocacy?				
	Is there				
Tell us more!	anything				
	else you'd like us to				
	know about?				
	Please tell				
	us!				
	us!				

You can put your name here if you like 😊

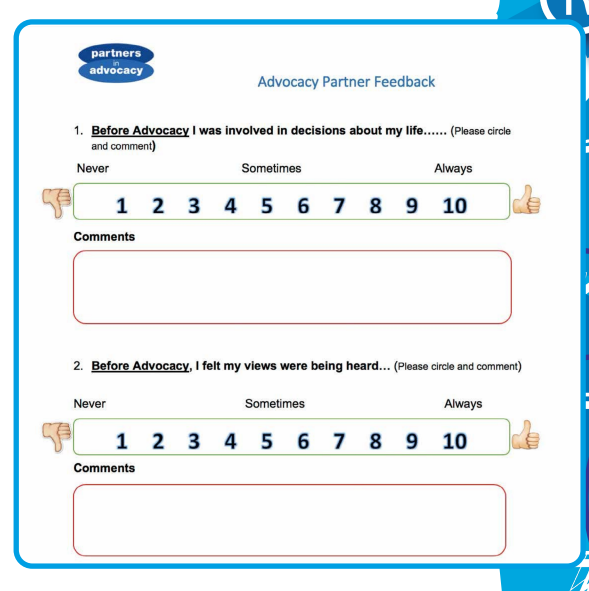
THANK YOU!!



While it is preferable to collect baseline information at the start, if that isn't possible – for whatever reason – it is worth asking a partner to reflect on their experience at the end of the partnership. Again, each independent advocacy organisation will need to make judgements about what is the best approach for its staff and client group.

The other possibility that it illustrates is that of embedding rating scales (see page 65) within questionnaires.

When visiting independent advocacy organisations, we were shown a range of questionnaires. Questionnaires can be any length, and presented in a range of styles. The important feature is, of course, the questions.



We have assembled a range of potential questions for different stakeholder groups – to inspire you to think about both open and closed questions that might be useful with different stakeholder groups. As advocacy partners are the focus of the examples provided, we have not included questions for them.

Questions for Collective Advocacy Members

One thing to keep in mind when measuring the impact of collective advocacy is that the outcomes are likely to be a mix of changes in the organisations or systems they are working to influence – and also changes for members, *individually*.

While collective advocacy groups in Scotland have some evidence of successfully influencing the way that services are offered, there is perhaps less recorded about how being a member of a collective advocacy group changes the individual members.

It is also likely that 'softer' impacts on services and the professionals who work in them go unrecorded. For example, changing attitudes to specific groups (e.g. elderly people, people who hear voices, etc), thinking differently, and even using different behaviours. Again, only by asking for feedback will independent advocacy organisations understand the impacts they have, and outcomes they achieve. It may be worth collective advocacy groups looking at the questions suggested for other professionals (see page 60) for inspiration.

We held a small focus group (a method that is discussed on page 64) with people who have been involved in collective advocacy, asking about the impact for them, personally – and heard some very rich stories.



"[Collective advocacy] has made a huge, positive difference to me – to learn about myself, and others. We are all there for the same reason – which helps to build trust. Some of the barriers I might experience socially are gone – because we are all there for same reason."

"[Collective advocacy is] a reason to get out of bed – a big part of my recovery. Some structure, something to do – that also might help someone else. Small steps that have helped me get back on my feet."

"We have had similar experiences, so we feel more equal...power dynamics are different....

Taking action, rather than passively taking what you are given...that is motivating!"

"I have learned more compassion, and I want to model that for others...so people can recognise the value of it. Kindness to yourself, and mostly other people."

"Meeting real people, who are having real feelings...our current news is so terrible, that it is kind of dehumanising. Collective advocacy is the opposite."

Some of the outcomes described in these statements include, increased social confidence, a sense of empowerment, a positive impact on recovery and better selfcare. Asking the right questions will make this evidence is available to organisations, to demonstrate some of the less obvious impacts of collective advocacy.



Useful questions for collective advocacy members

What do you believe has changed, for you, as a result of being a collective advocacy member?

Has being a collective advocacy member had an impact on other areas of your life? Y/N If yes, please describe.

How is being a member of a collective advocacy group different, in your experience, from other groups you may be involved with?

Has the experience of being part of a collective advocacy group resulted in your being more engaged in your community (joining other campaigns, being more aware of politics and power issues locally, etc.). Y/N If yes, please describe.

Would you recommend joining a collective advocacy group to others? Y/N If yes, why?

If you were invited to describe the value of collective advocacy to people who make decisions about whether or not to fund it, what would you want them to know?

Questions for family and friends

None of the independent advocacy organisations that we visited are routinely gathering feedback from family and friends of advocacy partners/collective group members. We understand that some may consider this an intrusion into the privacy of the partner/group member – and respect that this may be a principle that organisations are not willing to consider revisiting.



That said, there may be others – particularly those working with children and young people – who wish to consider this potential source of evidence.

One of the reasons it may be worth considering is that the people who are close to a partner/group member, but outside of their involvement with independent advocacy, offer a unique perspective on the difference that it can make.

It may well be that family/friends are a group that your organisations 'sample' on occasion, or you engage with intermittently – when opportunities arise, and it seems appropriate to ask the advocacy partner/collective member for permission to ask their family/friend for feedback. When asking permission, it is important to share the questions that you would like to ask. Remember to be clear about *why* the information is being gathered, and *how* it will be used.

Useful questions for family and friends

How would you describe the impact of being involved with independent advocacy for your friend/family member?

If they have spoken about the experience, what have they described?

Do you notice any changes in their behaviour or attitude, that you believe is a result of being involved with independent advocacy?

Would you encourage them to use independent advocacy again, in the future? Y/N

Why?

As a result of your friend/family members experience using independent advocacy, would you consider it for yourself in the future? Y/N

Why?



Questions for other professionals, who have experienced the advocacy partnership/collective group

Professionals who have experienced the difference that independent advocacy can make are very useful to hear from. Their feedback on the impact that independent advocacy can demonstrate its influence on services, and the professionals who deliver them.

Engaging with independent advocacy might mean that professionals begin to think, and even act, in new ways – because advocacy has shown them a way of making decisions that can meaningfully involve the person/people who will be affected by them.

If you never ask, you will never know about this wider impact.

Like friends and families, other professionals may be a group that you 'sample', or engage with when an opportunity arises – because of the relationship built through advocacy.

Because the questions you are likely to ask are about their own experience of advocacy, it isn't necessary to ask a partner/collective group member's permission.

Useful questions for professionals

Was working with advocacy involvement noticeably different to working without it? Y/N

In what ways?

Do you believe that advocacy had an effect on the choices that you were responsible for making, as a professional? Y/N In what ways?

Have you discussed your experience of independent advocacy with other colleagues? Y/N

If yes, what have you said about the experience?

Has your experience of advocacy affected your thinking, or the behaviours that you use, as a





professional? Y/N If yes, please explain.

Would you be likely to recommend independent advocacy to someone in the future? Y/N If yes, to whom – and why?

Useful technology

Most of the independent advocacy organisations we spoke with are using paper questionnaires, often delivered through the post, with enclosed return envelopes.

Most of these get fairly poor returns – and have therefore spent money unnecessarily.

We heard from those organisations who are using technology, that they have a significantly higher return rate. This is particularly true when the advocate is able to offer the questionnaire in person, using their phone or a tablet. It is also possible to send a link via email to an online questionnaire – which we heard also gets better returns than paper sent through the post.

As has been said already in this toolkit, it is for each independent advocacy organisation to consider the options, and create an approach to gathering evidence that suits their client group, and their available resources.

An additional benefit of using technology is that the programmes produce a report on the data that is input – so no time is required for this important task. The programmes will produce a report on the data, which the organisation can then analyse, and easily include in any reports.

The two programmes that we heard being used by independent advocacy organisations – in addition to specific database/case management systems that they are using – are <u>Survey Monkey</u> and Alchemer.



Survey Monkey is a free tool, while Alchemer requires a subscription, and offers a special charity rate. It is capable of more sophisticated collection/reporting of data and can be integrated within other on-line organisational systems (see website for further information). While Alchemer is a paid-for service, it might not cost an organisation too much more than it is spending on postage for evaluations already; though it is likely to produce much better returns on that investment, and Survey Monkey, are able to gather and report both quantitative and qualitative evidence, and both are relatively easy to use.

Microsoft Forms is another Survey tool that is often used and part of the Office 365 package which charities can access for free through a charity subscription <u>Microsoft Forms - Free tool to create online surveys, forms, polls, and quizzes (office.com).</u>

Individual interviews

Recommended for use with: Anyone

Interviews are an excellent way to gather evidence from the full range of sources mentioned in the section on questionnaires. In one sense, they are simply an alternative method for completing a questionnaire – with the advantage of being able to ask supplementary questions.

An interview presents an opportunity to gather a range of information – evidence of impact/outcomes, specific feedback on the way the service was delivered, ideas about how service delivery could be improved as well as specifics about methods or interventions that someone believes were important for them.

All of the questions suggested in the previous section on questionnaires would also be good interview questions.



The main drawback of interviewing is that it is a time-consuming means of gathering evidence. For that reason, it is likely to be used to supplement evidence gathered more consistently, and by other means. It is likely that conducting interviews is something that an organisation uses more infrequently, to supplement routinely gathered evidence, or to answer specific questions more fully.

For instance, arranging telephone interviews with a small number of people whose advocacy partnerships/groups have closed, to hear about their experience in more depth than a written questionnaire allows. Or arranging a small number of interviews with people with a shared characteristic which is under-represented amongst advocacy partners, enabling the organisation to learn more about how to be more accessible to this section of the population.

Interviews can be done in person, or on the telephone – and you may get the best response if you offer people a choice. For a questionnaire that has between 5-8 questions, with a number of qualitative questions being asked (hearing these stories is what makes interviewing worth the investment of time), you are probably asking people to participate in an interview that lasts 20-30 minutes.

You will want to think about how you will gather the evidence – make a record of the interview. Having a second person present to take a note is perhaps the easiest way to record an interview – or a voice recording. If you are making a voice recording, it is important to have clear permission from the person being interviewed, and assure them that the recording will be destroyed after it has been transcribed.



Focus groups

Recommended for use with: Anyone

Focus groups – or group interviews – offer something different to individual interviews, in that participants are often in conversation amongst themselves, so encouraged to think about different aspects of their independent advocacy experience than they might if only speaking to an interviewer.

The key is to invite/select people who have something in common – their advocacy issue or setting, a shared life feature or condition, etc. They need something that gives them a sense of connection – as well as making participants feel more at ease knowing that they have something in common, it also will help keep the conversation more focused than it might be otherwise.

How to run a focus group

Ideally there are 4-8 participants in a group – start with a small group, if it is your first time.

A focus group is not an ordinary conversation, as it follows a schedule of questions and is recorded in some way. Thought needs to be given to how the group will be recorded. The focus group can be voice recorded (be sure to get permission from participants) although it might be difficult to hear people talking over each other. It is hard to take notes and run a focus group at the same time – which makes the best option to have a second person who is taking a note of what people say. This could be a detailed note, or simply putting the main points on a flipchart.

Focus groups have the advantage that people who take part can listen and respond to each other as well as to the facilitator, meaning that the discussion can take new and interesting directions. People generally become quite energised when sharing their stories with one



another, so the facilitator will want to be careful not to let things go too far off track.

It can be useful to send questions out to participants in advance, so they come with thoughts to share. Just as with a questionnaire, you can combine questions relating to the outcomes you need to evidence, and also to get feedback about participants' experience of your organisation, and its activities. There are a range of useful examples in the section on questionnaires (page 50).

Focus groups can be carried out in a variety of settings, including the organisation's own premises or a local community venue. Ideally the area will be quiet, comfortable and reasonably private. Unless you are inviting other professionals, it is good practice to offer to cover travel expenses.

Rating scales

Recommended for use with: Advocacy Partners and Collective Group Members

Rating scales were singled out for their usefulness in the first section of the toolkit, particularly for use with Advocacy Partners and Collective Group Members. What makes them particularly useful is the fact that they were created to 'measure' change in a systematic and clear way. This makes them useful both for the organisation's learning, and also enabling the partner/group member to reflect on, and thus deepen, their own understanding of what participating in independent advocacy means for them.

Another thing that is useful about rating scales is their inherent flexibility – they can function, and look like, whatever the organisation believes will best suit its purpose for collecting evidence.



They can be a traditional numerical scale, a selection of images, an abacus (wooden counting frame), a drawing of a pizza – for which toppings represent intended outcomes. Or a real pizza for which toppings represent intended outcomes!

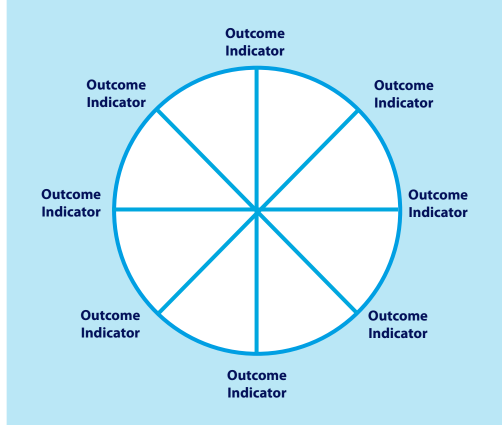
There are useful examples of paper-based scales available, and – on a paid for 'licensed' basis – including a range of support resources, at the social enterprise http://www.outcomesstar.org.uk.

We heard from a couple of the independent advocacy organisations that we visited, in preparation for writing this toolkit, that they are very pleased with what their use of the Outcomes Star – and that it is a worthwhile investment of just over £300 annually.

There is also a very good free resource available on the Evaluation Support Scotland website, called the Evaluation Wheel. This is, in essence, the same tool as an 'Outcomes star' – just visually different. The tool, and instructions for using it, can be found at http://www.evaluationsupportscotland.org.uk/resources/357.



Evaluation Wheel



Here is another version of an Evaluation Wheel.

While these wheels and the Outcomes Star look different, they are all simple 'rating scales':

- On which desired outcomes are identified (and placed on each 'spoke')
- On which an individual, or group, 'rate' themselves along a scale (places a mark on a point on each spoke, in relation to where they believe they are at present, in relation to the corresponding outcome.) It can be useful to ask the person doing the rating what that 'rate' indicates for them why have they selected it? This is important information, as it tells you what the rating actually means.
- The wheel/star is intended to be revisited at different intervals beginning and end, or other key stages so that a measurement (distance travelled) can be established.
- One independent advocacy organisation we spoke with said they had identified that if they asked for an Outcome Star to be completed in week six, they often got 'false data' as it was a common point for partner's optimism to dip. Organisations may initially wish to experiment with repeating the star/wheel (or other scales) at varying intervals in order to identify the best times to use it.
- Whether or not the advocate/group members wish to offer an opportunity for partners/one another to reflect on what the evidence gathered means to the person who filled it in – to describe what it says to them about their own journey – is a decision for each organisation.

This is the only resource in the toolkit that has this potential in-built.



An alternative rating scale: Pizza

Recommended for use with: Children and young people, people with a learning disability

Some advocacy partners/groups might engage more readily with a rating scale that is presented as something more familiar, and thus easier to engage with.

This scale works well in various formats, so can be easily adapted to different environments.

There are some settings – youth centres, various residential settings, community centres – where it may be possible to make an actual pizza. Alternatively, it is just as effective to create a 'wipeable' pizza base that is laminated (or even draw one on a flipchart/white board!)

What turns a pizza into a rating scale is creating a code for the toppings. For instance, pepperoni = confident to express my opinions, mushrooms = confident that I can influence decisions being made about me, onions = I understand my rights, cheese = feel I have power, etc.

The amount of each topping that is put on the pizza reflects the 'rating' that the partner/ group member is giving – making this type of scale slightly different. Rather than numbers on a scale signifying a position in relation to an outcome the amount of a representative topping signifies the position.

For instance – the 'baseline' pizza image might have very few toppings, or even none of certain toppings like cheese = I feel I have power, or onions = I understand my rights. Over time, the amount of toppings will likely increase. Therefore, it is still possible to make valid comparisons – noticing any increases.

Once a pizza has been completed, take a photo of it. This then becomes the record – against which future pizzas can be compared!



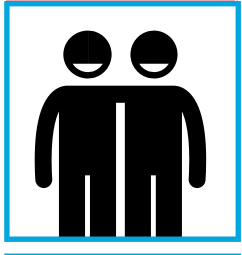
An alternative rating scale: Image-scale

Recommended for use with: Anyone



















Some advocacy partners and group members might be more comfortable relating to images than words. For that reason, this image-scale tool is included.

The images on the page depict people in a range of feeling states – which can be 'labelled' by the person who is using them as a rating scale. Like the evaluation wheel, start by being clear about the outcome you are asking them to 'rate' themselves in relation to – and then ask them to select an image that represents how they feel in relation to it.

Ask them to describe what the image says to them – how would they describe the image? Let them know you will be taking brief notes about what they describe – which you will read back to them at the end, to be sure that they are happy with what you have recorded.

Just like the evaluation wheel/Outcome Star, this is a tool to revisit on occasion. When using ANY scale, you can ask people to say a bit about why they chose the point on the scale/image – what does it mean to them? This also means that you are collecting both qualitative and quantitative evidence, which you can report on.

Just like the other scales, you might use this to establish a 'rating' in relation to a number of outcomes – with the individual choosing a new image, and describing it, for each.



Hurrah wall

For use with: Anyone

It was mentioned in the first section of this toolkit that there is likely to be a lot of evidence available throughout any given week that goes unrecorded, and perhaps even unrecognised, as *evidence*. This will most likely be thank you notes, text messages and emails.

Angus Independent Advocacy described the positive impacts that using a large white board in their reception area as a 'Hurrah Wall' has had for staff, and the organisation as a whole. Thank you notes and other forms of positive feedback are gathered and displayed on the wall – greeting everyone who comes through the door with evidence of the positive impact of independent advocacy in Angus.

In addition to offering a motivating reminder, every day, of the organisation's purpose and impact a Hurrah Wall is also excellent 'PR' – as most people sitting in the reception area will use the opportunity to look at it. This will include people who are considering using independent advocacy for the first time, current partners/group members, other professionals, and family/friends of partners.

It is good to clear the wall once a quarter, and at that time to consider the cards and notes as potential evidence of outcomes. Simply write each outcome on a sheet of paper, and put it on the floor – lined up in a row. Then read each note/card and – if appropriate – put it down beneath the outcome that you consider it to be evidence of.

There will be a number of things that won't actually be evidence of outcomes – simply notes of gratitude. That is fine – simply put the evidence you find in its appropriate column and then record it.



Observation, case notes and feedback

Apart from the Hurrah Wall, tools and approaches in this section of the toolkit are examples of how to structure requests for feedback, and methods for gathering it.

In addition to this type of evidence, advocates' own observations and case notes, as well as informal feedback provided by partners or group members, *are evidence*.

Generally speaking, it is best to consider observation and case notes as sources of evidence that substantiate – support the truth of – evidence provided by others, rather than as a primary source.

For instance, an **observation** that an advocacy partner appears to be using better self-care can be used to substantiate the partner's description of independent advocacy having a positive impact on their mental health.

The observation would not, on its own, constitute evidence of improved mental health – but it can be used to substantiate evidence that was provided by the advocacy partner/group member.

The same would be true for **case notes** – not to be considered a *primary source* of evidence, but a good secondary source which can be used to substantiate other evidence.

Informal **feedback** – that which hasn't been invited – from any of the stakeholder groups mentioned in this section of the toolkit can also be considered evidence. As described on page 33 all that is required is to ask permission to write down what has been said, and label it evidence.



Reporting on your evidence

Often when you report your evidence you will be reporting to *funders* – but they are only one of your potential audiences.

Current and potential partners/collective group members, related organisations, your Board, potential volunteers, friends and family of partners and group members are *all* potentially interested in your evidence.

Annual reports, your website and social media, promotional materials, volunteer recruitment, funding applications and fundraising campaigns are all opportunities to use the evidence you have gathered.

Remember to include both quantitative and qualitative data when sharing your evidence – for all audiences. Present quantitative data through visual representations such as tables, pie charts or graphs – as they can depict a lot of data in a focused and clear style.

Qualitative data is best presented in quotes or case studies. When writing case studies, it is best to use people's own words as much as possible. Ideally, they might be asked to write it themselves, or with some support.

Funders often indicate what they would like to read in reports, and in some cases may have stipulated the use of specific reporting templates. If no reporting template has been provided, be sure to provide evidence which is directly related to the outcomes/measures that you detailed in your funding agreement.

Enjoy sharing the evidence you gather of your organisation's outcomes in as many ways, with as many audiences, as is useful.



The SIAA looks forward to the opportunity to share evidence of members achieving the national outcomes for independent advocacy in Scotland – and knowing that there is an evidence base for the difference that independent advocacy can make.



Useful links and resources

National Development Team for Inclusion (NDTi) - Advocacy outcomes framework (ndti.org.uk)

The Lasting Difference - guide for sustaining non-profit organisations

<u>Advocacy: Models and effectiveness document from Iriss.</u> The <u>Iriss website</u> also has information on mental health, advocacy, and collaboration and involvement in social services.

Evaluation Support Scotland (ESS) - General evaluation information and support

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Partners in Advocacy

Dunfermline Advocacy

AdvoCard

Lomond and Argyll Advocacy Service

Advocating Together

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