

NSW Restrictive Practices Authorisation System User Guide

Part 4: Recording Outcomes and

Decisions

Version 5.0

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1. Purpose of the User Guide

1.1 Introduction

The Restrictive Practices Authorisation (RPA) User Guide (the User Guide) has been developed to assist NDIS Registered Service Providers (Service Providers) and Behaviour Support Practitioners (Practitioners) navigate the NSW (DCJ) RPA System (the System) in order to implement and comply with the RPA Policy and Procedural Guide. This user guide aims to provide the link between policy and practice.

The NSW Restrictive Practices Authorisation System User Guide *Part 4: Recording Outcomes and Decisions* takes users through the Outcome Summary and details how to record decisions made by the Restrictive Practices Authorisation Panel.

The User Guide has been updated and divided into 6 parts to allow users easier access to information. Each part is related to specific steps within the RPA System, therefore *Part 4: Recording Outcomes and Decisions* should be read in conjunction with all 6 parts (see following page).

This User Guide will be progressively updated as additional functions are added to the System.

Throughout the User Guide, **BEST PRACTICE SUGGESTIONS** have been included that your organisation may wish to use when establishing RPA processes.

Quick Reference Guides

Quick Reference Guides are highlighted throughout the guide. They can be found in the System under the Help menu.

1.2 How to use the User Guide

The RPA System User Guide has been divided into the following Parts:

RPA SYSTEM USER GUIDE



1.4 Authorisation Process Map



2. The Outcome Summary explained

2.1 Status of the Outcome Summary

The 'RPA with Outcomes' (*Outcome Summary*) is created when the release button is clicked on the RPA Submission Form.

→ Refer to User Guide Part 2: Submitting an Application and User Guide Part 3: Lodging a Submission

The status of each section of the Outcome Summary will be referred to through out this User Guide:



1	Status: New	Entering information into Sections 1 - 6 of the RPA with Outcomes is the first stage of recording the decision made at the RPA Panel. The Status of the RPA with Outcomes will be <i>New</i> .		
2	Status:	Clicking on ' <i>Release</i> ' after all information has been entered into Sections 1 - 6, opens up Sections 7 - 8 of the RPA with Outcomes.	Panel Annroval	
2	Pending Panel Approval	The status of the RPA with Outcomes will change from <i>New</i> to <i>Pending Panel Approval.</i>	& Consent can occur	
	Status:	Once all Panel members have approved the RPA with Outcomes, Consent is to be entered.	concurrently.	
3	Pending Consent	The <i>Complete</i> button is to be clicked which finalises the end to end RPA process.		
		The status will change from Pending Consent to Completed.		

2.2 Button Descriptors

The below buttons appear at the bottom of the screen during the different stages of the RPA with Outcomes:

Release	 This button releases the information to all the Panel members for approval. The information already entered in the RPA with Outcomes will be locked.
Back	• Clicking on this button will take you back to the RPA with Outcomes page.
Unsubmit	 Once all the information has been released, the information in the RPA with Outcomes is locked. Clicking Unsubmit unlocks the previous section and allows the applicant to edit information, if required. Once the RPA Form is released again, new notifications are sent to the Endorsing Practitioner and all Panel Members.
Save	• This button saves all information entered in the RPA with Outcomes.
Create Review	• Once the RPA with Outcomes is completed, a Review can be created if required.
Print	• This button opens up a PDF version of the RPA with Outcomes that can be printed or saved. This is what is sent to the NDIS Commission.

3. Convene the Panel

The implementing Service Provider is required to hold a properly constituted RPA Panel meeting which meets membership requirements.

3.1 RPA Panel for Planned Authorisation

The Panel can meet face to face, via teleconference, or webinar. The objective is for the Panel to come together to discuss the RPA application and make an informed decision as to whether the RPA submission will be approved, approved with conditions, or not approved.

- The Panel is required to evaluate the appropriateness of a proposed RRP and determine whether it is the least restrictive option available, or whether it will achieve the intended therapeutic outcome. The Panel will do this by considering the BSP, other relevant documentation and recommendations.
- The Panel is to have a regular schedule to enable:
 - 1. Systematic consideration and progression of applications.
 - 2. Regular monitoring, review and reporting of RRPs in accordance with the requirements set out by the NDIS Commission.
- The decision regarding authorisation must be recorded using the Outcome Summary.

No member of the Panel can also bring forward an application for the Panel's consideration.

Refer to User Guide Part 3: Endorsement and Assigning Panel Members for clarification of the roles required on a RPA Panel

> The Panel's decision to authorise a RRP is to be unanimous.

Unauthorised Restrictive Practices are Reportable Incidents.

3.2 Interim Authorisation, no Panel required

In exceptional circumstances, RRP may need to be rapidly implemented. Interim authorisation should be sought as soon as practicable, not exceeding 1 month.

A Senior Manager of the implementing Service Provider can provide interim authorisation based on an interim BSP written by a behaviour support practitioner.

The Senior Manager should consider the content of the interim behaviour supports and be satisfied that the strategies outlined represent the least restrictive option.

- The decision regarding authorisation must be recorded using the Outcome Summary.
- Interim Authorisation should be for the shortest duration required to minimise the risk.

Full (planned) authorisation should be obtained within 6 months, or the restrictive practice(s) should be discontinued. Application for an Interim RPA follows the same process as a Planned RPA until the assignment of Panel Members.

→ Refer to User Guide Part 2: Starting a Submission and User Guide Part 3: Lodging a Submission

Refer to User Guide Part 3: Endorsement and Assigning Panel Members for clarification of the roles required on a RPA Panel

4. Outcome Summary

Outcome Status: New

Before you can start a RPA with Outcomes in the RPA System, a RPA Submission must have been entered in the System and released.

→ User Guide Part 2: Submitting and Application and User Guide Part 3: Endorsement and Assigning Panel Members.



Having access to the System during the Panel means information can be immediately recorded on the Outcome Summary.

<u>Note</u>: the session will timeout if left inactive for 10 minutes. If this happens, log back in.

4.2 How to start an Outcome Summary

Once the Panel has discussed the application and have come to a unanimous decision, the Senior Manager enters the Panel decision into the System.

Section 1 of the RPA with Outcomes

1. NDIS Participant Details

This section has been pre-populated from the information entered into the RPA Submission Form.

To change or update participant details in the System:



→ Refer to User Guide Part 2: Submitting and Application

Section 2 of the RPA with Outcomes

2. RPA Application Details

Section 2 has been pre-populated with the following:

- Submission ID number,
- Behaviour Support ID (if applicable), and
- Submission Type
- Behaviour Support Plan Expiry Date

To open the RPA Submission press Ctrl and click on the Submission ID hyperlink. The RPA Submission will open in a new browser tab.

Information entered in the RPA Submission cannot be changed via the Outcome.

Outcome Status: New

Outcome ID Number

Each RPA Form will have its own unique Outcome ID number.

This number can be used when searching within the System, and is linked to the Submission ID number.

Outcome Status: New

Section 3 of the RPA with Outcomes

3. Proposed Restrictive Practice

This section has been pre-populated with the following information from the RPA Submission Form:

- Service Setting
- Behaviour of Concern
- Restrictive Practices Category
- Description of Proposed Practice/Strategy
- Fade-out strategies



The Proposed Restrictive Category information entered in Section 6 of the RPA Submission

Form can be viewed by clicking the View Details button.

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Hover Help is available throughout the RPA System. Clicking on the '?' symbol, will open up a brief summary that may provide assistance to you when entering information in the System. The Proposed Restrictive Practices information will open in a pop up box.

To close the Restrictive Practices pop up box, click on the *back* button.

Quick Reference Guides

Restrictive Practices Authorisation— Complete a RPA Outcome Summary

4.3 Entering the RPA Outcome Decision

Section 4 of the RPA with Outcomes Form

4. RPA Outcome Decision

	Chose the decision the RP	A Panel has made from the drop down list:	restrictive practice is to be unanimous.
Step 1	Decision		
	Please select 🔹		
	Please sel Approved Approved Not Appro	ect – with Conditions oved	
Step 2	Enter the Decision Date:	Decision Date	
Step 3	Enter the Meeting Date:	Meeting Date	The Panel needs to discuss the restrictive practice and come to a decision. Therefore, the
	1		Decision Date entered cannot be before the

Outcome Status: New

The Panel's

authorise a

Meeting Date.

A description of why the authorisation was given/declined must be recorded for each RPA submission to a Panel.

Recording what was discussed and why a RPA Panel made a particular decision in the 'Why was the authorisation given/declined' field will assist Service Providers to implement, monitor and report on their Organisation's authorised practices.

Step 4



When recording the Reason for Decision, remember that most recipients of the Outcome Summary will not have been present at the discussion, so it is important to provide a detailed summary of the Panel's discussion and rationale for the decision.

The 'Conditions' field is an optional field. If the RPA Panel has applied specific conditionsStep 5 on the authorisation of a practice then this is where they should be recorded. RPA Panels should schedule a Review to monitor actions taken regarding the conditions.

Approved - with Conditions

If the decision of the RPA Panel is *Approved with Conditions*, the Conditions field becomes mandatory.

When Approved - with Conditions has not been selected, the Conditions field is not mandatory.

 Step 6
 In the 'Panel Recommendations' field, any additional suggestions/actions that the Panel have determined would be useful for the situation presented in the submission should be recorded here. These are not conditions and the field is optional.
 RPA Expiry Date

 Step 7
 Enter the RPA Expiry Date:
 RPA Expiry Date

 The RPA Expiry Date cannot be prior to the Panel Meeting Date.
 The RPA Expiry Date cannot be prior to the Panel Meeting Date.

Refer to Section 2 of User Guide Part 1: Accessing the System Interim authorisation: Cannot exceed a *total* of 6 months from the decision date.

Planned authorisation: Cannot exceed a total of 12 months from the decision date. **Consent** Consent cannot expire before the RPA Expiry Date.

Outcome

Status: New

	If a Review is required as part of the lifecycle of this particular authorised restrictive pratice, then the 'Panel requires a review of use' box should be ticked:	Outcom Status: N
Step 8	Panel requires a review of use	
	Once the box has been ticked a review date should be entered in the 'Review to occur b field.	ру'
	The Review to Occur by date is not the same as the RPA Expiry Date.	
	The Review to Occur by date should occur at an interval/s during the authorisaiton peri and not shortly prior to the RPA Expiry Date.	od
	Where a review date is set by the Panel, the System will automatically generate a draft. If a review is not required, do not tick the box.	
Step 9	If the Panel decides that a review is required within the authorisation lifecycle, then ent the 'Review to Occur By' Date: Review to occur by	ter Prior to a RPA Expiry Date, a new RPA Submission Form is to be started.
	The 'Review to Occur By' date is <i>not the same</i> as the RPA Expiry Date. The 'Review to Occur By' date should be <i>prior to</i> the RPA Expiry Date.	A Review is a review of a current authorisation, it is <i>not</i> re-authorisation.
	Where a review date is set by the Panel, the System will automatically generate a <i>draft</i> Review Submission in the System.	For further detail
	Reviews	refer to
	A Review can also be initiated after the RPA with Outcomes has been completed, if required due to a number of	User Guide Part 5: Reviewing Implementation
	circumstances including if a practice is to	Progress

circumstances, including if a practice is to be ceased or has been superseded.

4. Outcome Summary

Section 5 of the RPA with Outcomes

5. Add any additional documents to support the decision

The Panel can upload any notes or other supporting documents from the Panel meeting into the System, as required, by choosing the Add Files button.

Section 6 of the RPA with Outcomes

6. Documents Required for Next Review

If a Next Review Date has been entered by the Panel, the Panel can choose to request additional documents be provided at that Review. This will be recorded within the System, and will be mandatory to provide at the first Review (but not for any adhoc reviews).



This is not a mandatory field within the System. If additional documents are not requested by the Panel, the Panel does not have to complete this field within the System.

Once all information has been added to the RPA with Outcome, click the

Release button.

The status of the RPA with Outcomes will change from New to Pending Panel Approval.



 \searrow All Panel Members will receive an email notification requesting they log into the system and Approve

the RPA with Outcomes.

Refer to **User Guide Part** 5: Reviewing Implementation **Progress**

Section 7 of the RPA with Outcomes

7. Evidence of Formal Consent

Enter the Consent Giver details in the System:

- Enter the Date of Consent and the Consent Expiry Date.
- Provide a description of the type of consent provided. For example, verbal or written consent.
- The Senior Manager can upload consent documents to the System by choosing the Add File button, if required.
- Once all consent details are entered into the System, either:
 - I. click the Save button, if you are still waiting for the Panel to approve, or
 - II. Click the **Complete** button if all Panel members have approved the RPA with Outcomes.

Consent is a mandatory requirement for the use of a restrictive practice.



If Panel Members have approved the RPA with Outcomes prior to Evidence of Formal Consent being entered in the System, the status will change from *Pending Panel Approval* to *Pending Consent*. Outcome Status: Pending Panel Approval

Section 7 and Section 8 can occur concurrently, and the Outcome Status will reflect the current status.

Reference Documents:

Appendix 3: Consent.

4.4 Panel Member Approval

Section 8 of the RPA with Outcomes

8. Panel Member Approval Once the RPA with Outcomes has been *released*, all Panel Members will receive an email notification stating that there is one or more outcome summary(s) awaiting endorsement on the System. All Panel Members are required to log into the System within 24 hours, to Approve the Outcome Summary. Approving the Outcome Summary indicates that the Panel Members agree with the decision and Reason for Decisions recorded in the Outcome Summary. The status will then change from *Pending Panel Approval* to *Pending Consent*.





Outcome Status: Completed

4.5 Completing the RPA with Outcomes

Once consent details and all Panel approvals have been entered in the System, the Senior Manager completes the

RPA with Outcomes by clicking the **Complete** button.



The status will change from *Pending Consent* to *Completed*.

Once completed, the System will automatically send a notification email to the RPA Submission applicant to advise that acknowledgement from all the Panel Members has been received, and the decision for one or more RPAs (Submission ID numbers) has been finalised.

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The Service Provider can now **Print** the Outcome Summary form to provide to the NDIS Commission to meet their legislative requirements.

Outcome Summary is ready to view

When approval from all Panel Members has been received, the System will send a notification to all involved in the RPA Submission advising that the Outcome Summary is complete and ready to view. Subject: NSW Restrictive Practices Authorisation - Outcome Summary Awaiting Approval Dear Panel Member.

You have the outcome summary [Outcome Summary ID number] awaiting your endorsement on the NSW restrictive practice authorisation (RPA) portal.

The submission is for [Participant name], [DOB] and the panel held on [date of Panel] convened by [Panel convenors name] on the Panel.

Please provide your endorsement of the decision by [Date by which endorsement is required]. You can access this application and all associated material at https://rpa.facs.nsw.gov.au

Should you have any questions regarding this application, please contact the panel convenor, [Panel convenors name] at [Panel convenors email] or [Panel convenors phone number]





Appendix 2: Documents applicable for an RPA Submission

the following: Documents considered evidence for the use of RPA Panel Members include, but are not limited to,

		• •	
	Behaviour Support plan	Consent Interim	Interim RPA Submissions
•	•	• •	
One Page Profile	Plan Functional Behavioural Analysis	Consent Behaviour Support	Planned RPA Submissions
• •	• •	•	Q
Behaviour Assessment Report PRN Protocol	Evidence of Implementation Training Lifestyle Plan	Data Collection Summary	ther documents you could RPA Submi
	• •	• •	consi issior
	Co Ris	<u> </u>	ider in ns

Environmental Restraint • N	Physical Restraint • Ru N	Mechanical Restraint N	Chemical Restraint • P	Seclusion • Se
o additional documents are required	estraint Register (This is the same as the data you have submitted to the DIS Commission as a part of your reporting requirements)	estraint Register (This is the same as the data you have submitted to the DIS Commission as a part of your reporting requirements)	RN Protocol and/or Medication Chart 1edical Reports Iformation on side effects and symptoms of overdose	eclusion Register (This is the same as the data you have submitted to the DIS Commission as a part of your reporting requirements)

It is best practice to provide more than just the mandatory documents.

submission. Not enough detail in the submission could affect the RPA Outcome. You need to provide enough detail so Panel Members can make an informed decision about the

Appendix 3: Consent

Consent is the permission given by the person (where they have the capacity to consent), or the person with authority to consent on that person's behalf (where they do not have capacity to consent).

Consent must be voluntary, informed, specific and current. Consent requirements for RRPs are summarised in the below table: RPA Consent Requirements. *Refer to section 4.3 of the Restrictive Practices Authorisation Policy for further information.*

Consent is needed to use a RRP as a component of an overall BSP.

The Service Provider must ensure that the person, or their consent giver, has been informed of the use of RRPs; the requirement for the RRP to be submitted to the System; and the Outcome Summary from the Panel.

Consent must be obtained from the participant or their guardian at two different points within the System:

1. Submit a RRP in the System

- Ticking the box at this point indicates that the Consent Giver is aware that an RPA submission is underway.
- Ticking the box does not imply a formal consent to implement an authorised practice. The Service Provider must ensure that the Consent Giver has been informed this could be by the Practitioner or the Service Provider.

2. Consent to the Outcome Summary

• Providing evidence of Formal Consent at this point indicates that the Consent Giver has been informed of the outcome of the Panel. The panel convenor is required to complete this section and is responsible for ensuring the Consent Giver is notified of the outcome.

Adults (18 years or over)	Young people (16-18 years)	Children (under 18 years) subject to court order reallocating parental responsibility	Children (under 18 years) <i>not</i> subject to court order reallocating parental responsibility	Person
Either: a)The person where they have the capacity, or b)Guardian with a restrictive practices function	Either: a)The person where they have the capacity, or b)Guardian with a restrictive practices function	Person with parental responsibility+	Parent of Guardian*	Physical or Mechanical Restraint
Either: a) The person where they have the capacity b) The person responsible c) The Guardianship Division	Either: a) The person where they have the capacity b) The person responsible c) The Guardianship Division	Person with parental responsibility+	Parent of Guardian*	Prac Chemical Restraint
 Either: a) The person where the capacity, or b) Guardian with a restrictive practices function, or c) The RPA Panel mechanism± 	Either: a) The person where they have the capacity, or b)Guardian with a restrictive practices function, or c) The RPA Panel mechanism±	Person with parental responsibility+	Parent of Guardian*	tice Environmental Restraint
 Either: a) The person where they have the capacity, or b) Guardian with a restrictive practices function 	PROHIBITED	PROHIBITED	PROHIBITED	Seclusion

- * and Young Persons (Care and Protection) Regulation agency in accordance with Clause 26 of the Children With approval of the principal officer of the designated <u>+</u>
- $^+$ court order must be provided. reallocating parental responsibility, evidence of the For children who are subject to a court order

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2012 as appropriate.

 $\mid +$ consent in certain circumstances. access) strategy may be implemented in the absence of environmental restraint (e.g. response cost or restricted The RPA mechanism may direct that an authorised

Notes:

Guardian Division can consent. Medication (or dental) Treatment and can only the behaviour, while not psychotropic, fall under Special Androgen-reducing medications prescribed to control

establish and maintain a RPA mechanism which authority does not release the registered NDIS provider use of a regulated restrictive practice by its staff. evaluates, authorizes and monitors all instances of the from the ethical imperative to have access to or to The consent of the person(s) with appropriate legal