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## Insert the article title here

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Some abstract goes here

## 1. Introduction

As is common in cognitive psychology, most visual search literature has focused on how the average participant performs in the task, despite it being well known that there is a great deal of variability between one subject and the next. From Treisman's work on Feature Integration Theory [20] to the latest incarnation of the Guided Search Model [22], we now have a good understanding of what makes some search targets effortless to find, while others require careful inspection. However, these theories and models have neglected the question of why some observers find visual search so much harder than others. These differences can emerge from several different sources of variation: tiredness [12], information-processing ability, speed-accuracy trade-off, motivation, visual impairments [15], and search strategies [1]. Although their existence has previously been noted [12], a rigorous examination of individual differences in visual search has been largely ignored and questions about their importance and stability remain relatively under explored.

In the current study, we focus on one source of individual differences in visual search: strategy. By search strategies, we refer to a collection of behaviours that all observers can freely choose from when they search a display. Examples include choosing to adopt a systematic left-to-right and top-to-bottom strategy [6], or the type of guided search behaviour in which locations that are more likely to contain the target are prioritised [22].

A striking example of the effect of strategy is given by Boot and colleagues [1]. They asked participants to monitor a cluttered display for an object changing colour or object onset. Large individual differences were found with respect to the number of saccades participants made while monitoring the stimulus, and this was negatively correlated with detection performance. Proulx [17] demonstrated that the different weighting between top-down and bottom-up factors in a conjunction search task explained some of the variation in how participants did in an attentional capture task. However, only a third of participants were unaware of the search strategy they employed.

Eye movement strategies have also been shown to be an important source of individual differences in visual search tasks. Nowakowska, Clarke & Hunt [14] designed a simple search paradigm (referred to here as the Split-Half Search Array paradigm) to discriminate between optimal [13] and stochastic [3] search strategies involving arrays of line segments (see Figure 1). The line segments were arranged such that those on one side of the display all had a very similar orientation, while those on the other side had higher variance. This was done so that if the target appeared on the homogeneous side, it would be highly salient. Trials in which the target appeared on the heterogeneous side of the display would be much harder. With these stimuli, the optimal eye movement strategy is to only search the heterogeneous half of the stimulus, as targets on the homogeneous side can be detected with peripheral vision. While some participants initially searched the displays near optimally, others carried out strategies counter to this, failing to even match the performance of the stochastic searcher. Furthermore, the degree to which they made saccades in line with the optimal search strategy was strongly correlated with their reaction times. A related version of this paradigm has been used in research in (simulated) hemianopia [15,16]. . . .

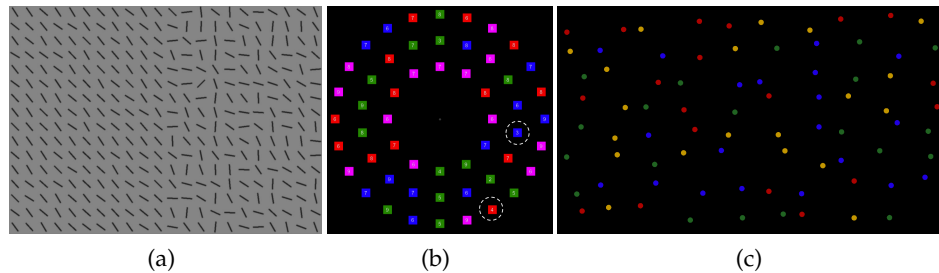
A similar range of strategies, from random to near-optimal has been found by Irons & Leber with the Adaptive Choice Visual Search (ACVS) paradigm [7], designed to explore how feature-based attention is used. This paradigm involves stimuli made up of small coloured boxes (red, blue, green and a fourth colour

that varies from red, through purple, to blue and back again) with numerals written inside them (See Figure 1). The target is defined as a red or blue box with a numeral between 1 and 4, and on each trial a target of both type is present. The participant's task is to find one of either target as quickly as possible and report the numeral. One trials in which the fourth box colour is red (or close to red), participants should search through the blue boxes and report the blue target, as there will be fewer distracters. As the fourth colour changes to through purple to blue, participants should update their strategy and search for the red target. The results showed that participants varied substantially along two key dimensions: how frequently they used the more effective target feature to search (varying from chance performance to near optimal), and how often they changed search features. Further work has shown that these differences are stable over time (between one and ten days) with test-retest correlations of around  $r \approx 0.8$  [8].

Another example of differences in search strategy comes from the foraging literature [10, 11]. In this context, foraging is a visual search task in which there are multiple targets on each trial. Participants were asked to search through a set of items from four categories, two of which were classed as targets. In the conjunction condition (i.e. searching for red-horizontal and green-vertical line segments among red-vertical and green-horizontal distracters), most observers searched in runs of one target category or another. This strategy has previously been observed in animal foraging literature [5]. However, a sub-set of observers, termed 'super-searchers' showed no switch cost.

Previous research has investigated the relationship between these behaviours to psychometric measures, but to date, these differences have not shown strong correlations with other attributes. In the ACVS paradigm, Irons & Leber found no evidence of a correlation between the proportion of optimal choices made by observers and measures of visual working memory or trait compulsivity [7]. Similarly, the differences foraging behaviour are not accounted for by working memory or inhibitory control [9].

A common theme emerging from the studies in the observation that individual strategies



**Figure 1.** Example stimulus from the (a) *split-half*, (b) *adaptive choice* and (c) *foraging* paradigms.

vary in their degree of effectiveness or optimality. However, “visual search” encompasses a wide range of specific tasks, each tapping into a different aspect of behaviour (e.g. feature-based attention, information sampling). The aim of the present study is to investigate the extent to which individual differences are stable across different visual search paradigms. Are observers who use the optimal strategy in the split-half search arrays also more optimal in the ACVS task? Does it make sense to talk about ‘super-searchers’ who show above average performance in a range of search tasks? Are the super-foragers consistently better or worse than more typical searchers in the other two paradigms? As a secondary question, we will measure the test-retest reliability of the differences found in the split-half array paradigm.

## 2. Methods

The methods and planned analysis for this study were registered on the Open Science Framework<sup>1</sup> before data collection started.

### (a) Participants

64 students from the University of Aberdeen took part in this study. Participants were compensated for their time with either course credit or £15. All participants will sign a form giving informed consent. The study was approved by the University of Aberdeen Psychology Ethics Committee.

Sample size was determined in part by a power analysis, and in part due to constraints with counter-balancing.  $n = 64$  participants means we should be able to detect correlations

<sup>1</sup>insert URL

with  $r > 0.34$  with  $\alpha = 0.05$ ,  $\beta = 0.80$  between the different visual search paradigms. This compares to the already established value of  $r \approx 0.8$  for the reliability for the ACVS paradigm.

### (b) Materials and Procedures

The study consists of three paradigms from the visual search literature in which large individual differences have been noted found [7,11,14]. Example stimuli can be seen in Figure 1. A brief overview of each paradigm is given below, with full details in *supplementary materials*. The three tasks will be completed over two sessions, approximately one week apart. The split-half array search will be run in both sessions allowing us to measure test-retest reliability. The order in which participants completed the tasks was counter-balanced. There are 16 different possible orders of tasks/conditions; we will run four participants in each order for a total of 64.

The display was presented on a 17-inch CRT monitor with a resolution of  $1400 \times 1050$ . Stimulus generation, presentation and data collection were controlled by MATLAB and the psychophysics and eyelink toolboxes [2,4] run on a Powermac.

#### (i) Split-half Array Search

Stimuli consisted of arrays of black oriented line segments against a grey background. Each line segment had a length of  $\approx 1.6^\circ$  of visual angle irrespective of screen resolution. The target was oriented  $45^\circ$  clockwise, while the distractor items had a random orientation with a mean of  $45^\circ$  anti-clockwise. The variance was low ( $18^\circ$ ) on one half of the display to create a homogeneous texture, and high ( $95^\circ$ ) on the

other side to create a heterogeneous texture. This means that when the target is present on the homogeneous side of the stimulus, it can be easily be detected with peripheral vision, but when it is in the heterogeneous half, it is much harder to detect. There were a total of 160 trials and homo- and heterogeneous sides of the display were randomly varied from trial to trial. The position of the dominant eye was recorded using a desktop-mounted EyeLink 1000 eye tracker (SR Research, Canada).

This paradigm was carried out twice to give us an estimate of how consistent participants are in their search strategy over time. The two sessions were identical.

## (ii) Adaptive Choice Visual Search

The ACVS was based on the task described in [7], Experiment 1. Each search display was composed of 54 small squares arranged in three concentric rings around fixation, with 12, 18 and 24 items in the inner, middle and outer rings respectively. The same screen was used as in the Split-half Array Search, however, due to changes of screen resolution, the size of the squares changed. Participants, were sat  $\approx 47$  cm from the screen. Of the 54 squares, 13 were red, 13 were blue, 14 were green and 14 were "variable". Variable distractors change colours from trial-to-trial according to a 24 trial cyclical pattern: the distractors would be red for 5 trials (called a "red plateau"), then across a period of 7 trials, they would change colour from almost red to magenta (at the fourth trial in the transition) to almost blue. The variable distractor would then be blue for 5 trials (blue plateau), and then transition back from almost blue through magenta to almost red.

A white digit appeared inside each square. Two targets - a red square and a blue square each with a digit between 2 and 5 - were embedded in every search display. The two target digits were always different, to enable us to distinguish the chosen target. The remaining red, blue and variable squares all contained digits between 6-9. Green squares could contain any digit between 2-9. The location of the targets and distractor within the search display were randomized on each trial.

Participants were informed that the search displays would contain two targets on every trial, that they need only find one target on each

trial and that they were always free to search for either one.

## (iii) Conjunction Foraging

The foraging task was based on [11] and [10]. Participants completed the feature foraging and conjunction foraging tasks on separate days, with the order counterbalanced (was it counterbalanced?).

In the feature foraging task, search displays contained 80 small circles, 20 red, 20 green, 20 blue and 20 yellow. Stimuli were arranged in a  $10 \times 8$  grid, but the position of each item within the grid space was jittered to create a more random spatial arrangement. The location of item colours to grid locations was completely randomized.

For half of the participants, targets were red and green circles, and for the other half of participants, targets were blue and yellow circles. Participants were asked to collect all of the targets within a trial by using the mouse to click on each target. Clicking on a target caused it to disappear from the display. If the participant clicked erroneously on a non-target, the trial was immediately ended and a replacement trial was begun.

In the conjunction foraging task, search displays were composed of both circles and squares. For half of the participants, the shapes were red and green (equal numbers of red circles, red squares, green circles and green squares), and for the remaining participants the shapes were blue and yellow. Targets were defined by conjunctions of colour and shape (e.g., red squares and green circles, with red circles and green squares as distractors). The assignment of targets and distractors was assigned at random for each participant. The procedure was otherwise identical to the feature foraging task.

# 3. Results

## (a) Replication of each paradigm

### (i) Split-half Array Search

Our results are in line with the original split-half array study [14]. Only trials with a correct target absent/present response were included in the analysis. Some participants performed exceptionally poorly and were removed from

further analysis (full details given in the supplementary materials). All reaction times were  $\log_2$  transformed. In order to characterise an individual's behaviour in this task, we will compute the proportion of the first  $n$  fixations that were on the heterogeneous (difficult) side of the stimuli, over all correct target absent trials. Previous work [14] demonstrated a strong correlation between this metric (for  $n = 5$ ) and reaction times ( $r = .53$ ).

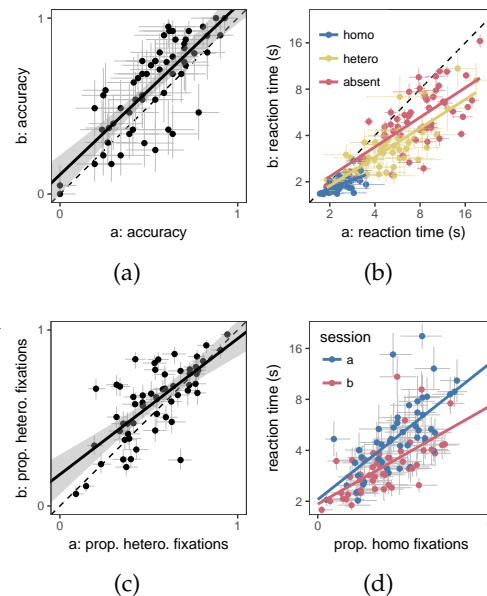
The correlation between accuracy and reaction times between the two sessions is shown in Figure 2(a, b). We can clearly see that there are large differences from one participant to the next in terms of both the proportion of hard targets found, and reaction times. Furthermore, test-retest reliability appears to be reasonable, with Pearson's  $r \in [0.65 - 0.86]$  (95% confidence interval) for accuracy in finding targets on the hard heterogeneous half of the display. We get similar scores for the correlation between sessions  $a$  and  $b$  for heterogeneous targets, ( $r \in [0.66 - 0.87]$ ), homogeneous targets ( $r \in [0.56 - 0.82]$ ) and target absent ( $r \in [0.65 - 0.86]$ ). The reduced correlation for the homogeneous targets is likely due to the restricted range.

We can also look at the initial search strategies adopted by our participants 2(c, d). Again, we see large and stable individual differences across the two sessions (test-retest  $r \in [0.57, 0.83]$  for the proportion of the first five saccades to the heterogeneous half of the display for target absent trials). More importantly, as with [14], we see that the search strategies give a good correlation with reaction times in both session  $a$ ,  $r \in [0.56, 0.84]$  and session  $b$ ,  $r \in [0.50, 0.81]$ .

## (ii) Adaptive Choice

Participants with accuracy more than 3 SD below the group mean were excluded from analyses. For RT analyses, trials with RTs less than 300ms or more than 3 SD about the participant's mean were excluded.

Two measures of individual strategy use were used: 1) Optimal choices, defined as percent of plateau trials in which the individual chose the optimal target (i.e., the target with the fewest distractors. When the variable distractor was red, the optimal choice was blue, and vice versa), and 2) Switch rate, the percent of trials



**Figure 2.** Each point represents a participant and the error-bars indicate 95% confidence intervals. Correlation between the two sessions of the *split-half* paradigm for (a) accuracy (TP-heterogeneous trials only); (b) reaction times and (c) search strategy (TA trials only). (d) Initial search strategy correlates with reaction times in both sessions.

in which the individual switched target colour (i.e., the colour chosen on trial  $N$  was different to the colour chosen on trial  $N-1$ ).

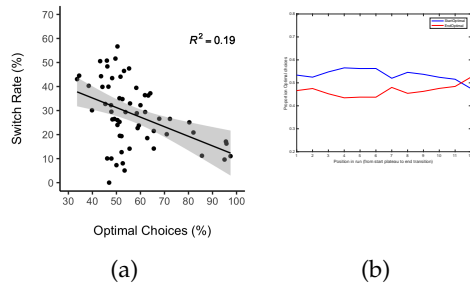
The results for the ACVS were consistent with previous findings [7,8] (see Figure 3). shows that there were wide individual differences in both percent option (range 33.62% - 97.50%,  $M = 56.99$ ,  $SD = 14.75$ ) and switch rate (range = 0% - 56.68%,  $M = 28.46$ ,  $SD = 13.54$ ).

## (iii) Conjunction Foraging

Only completed, accurate trials were analysed. RTs were defined across the entire trial (i.e., from the start of the trial until the final target was collected). The main measure of interest was average run length per trial. A run was defined as a succession of one or more of the same target type, which was followed and preceded by the other target or no target. The average run length was the average number of target selections in a run.

The multiple-target foraging results were also in line previous findings [10,11]. As expected, run lengths were shorter for feature





**Figure 3.** Needs checked and updated. replace png with pdf!

foraging ( $M = 3.24$ ,  $SD = 3.19$ ) than conjunction foraging ( $M = 10.64$ ,  $SD = 7.01$ ),  $t(55) = 8.03$ ,  $p < .001$  (PREFER DIFFERENT STATS HERE? CI [-9.25,-5.55]) suggesting more frequent foraging for multiple targets concurrently when those targets were defined by features than by conjunctions. However, both showed significant individual variation: Feature run length varied between 1.87 and 20, and conjunction run length varied between 1.00 and 19.67. Figure 4 depicts run length across the trials separately for each individual. Twenty “super-searchers” were revealed, identified by short run lengths for both feature and conjunction targets.

## (b) Correlations Between Paradigms

The results above demonstrate that we have successfully replicated the previous findings around individual differences in visual search strategy in each of the three paradigms. Furthermore, the split-half paradigm has a test-retest reliability of  $r \in [0.65, 0.86]$ , similar to that of the ACVS task ( $r \in [0.72, 0.9]$ ), suggesting that the strategies different individuals chose to use in each of these tasks are somewhat stable. Given this, we now investigate the extent to which an individual’s performance in one of the tasks tells us about how well they will do in the other two.

The results presented in Figure 5 convincingly demonstrate that there are no strong correlations

between the strategies by observers in split-half, adaptive choice and foraging visual search paradigms<sup>2</sup>.

While there is no indication of a relationship between strategies across the different paradigms, perhaps there is still a correlation between the reaction times due to some participants having higher levels of motivation than others, or stable speed-accuracy trade-offs between the different tasks. Differences relating to their information-processing speed and their ability to ignore distractors would also play in to such a correlation. However, as with search strategy, we again find all the correlations to be weak, typically  $0.27 < r < 0.30$  but positive (Figure 5). None of these correlations are statistically significant ( $p > 0.05$ ). Even if we optimistically take all the data together as suggesting a robust correlation in reaction times from paradigm to paradigm, the upper bound of  $r = 0.30$  means that this correlation only accounts for at best 10%<sup>3</sup> an individual’s performance.

## 4. Discussion

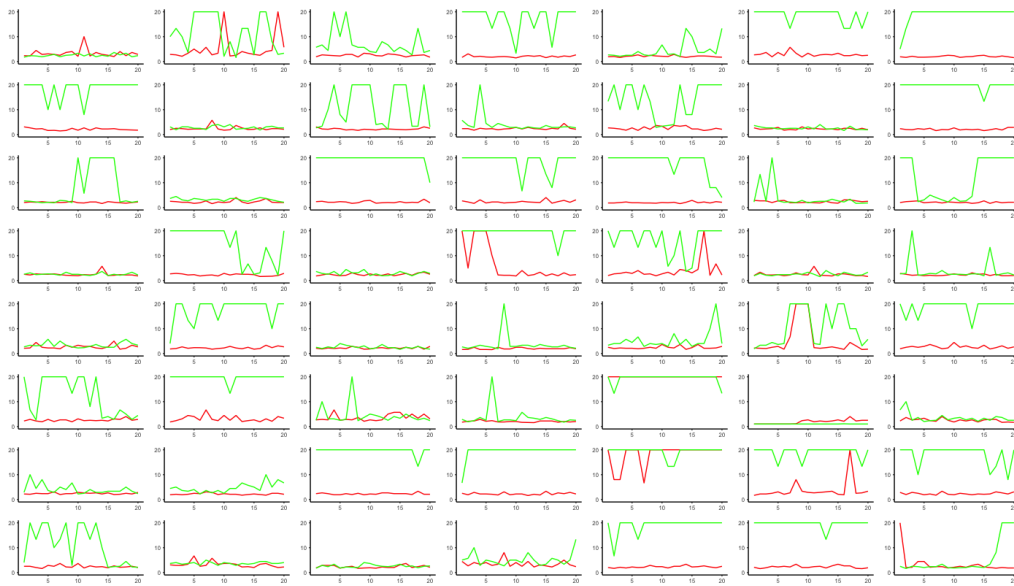
While we have successfully replicated the range of individual differences found in each of these studies, with a larger sample size than the original papers, the between-paradigm correlations give at most  $R^2 = 0.1$ , and this is a generous interpretation that would fail to pass the usual criteria for null hypothesis significance testing. Knowing how one person will behave in one of these paradigms apparently tells us very little about how they will perform in the others. This can be contrasted with the relatively high test-retest correlations of in the split-half paradigm and for the adaptive choice paradigm [8]. [AM I RIGHT IN THINKING ARNI AND CO HAVE MEASURES FOR THEIR PARADIGM???

I think he sees it as a positive thing that they tasks don’t correlate, because it suggests they capture unique variation in behaviour glass half full approach

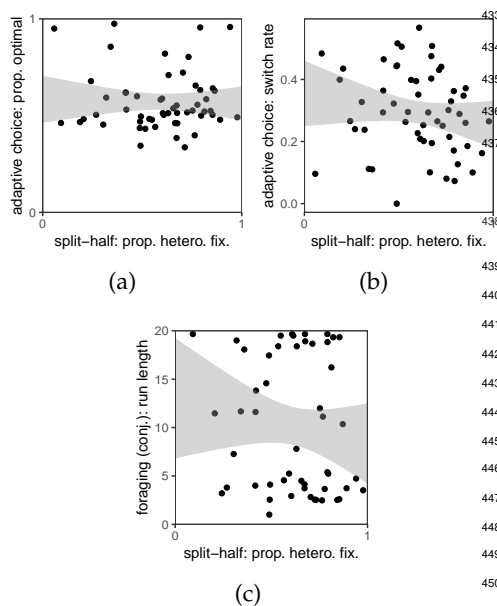
Mention [21]. From [17] - *Vogel and colleagues have found that an individual’s ability to remember a greater number of items using working memory is related to a filtering capacity in visual search that suppresses attentional capture by*

<sup>2</sup>More comparisons are presented in the supplementary materials.

<sup>3</sup>i.e.,  $R^2 = 0.3^2 = 0.09$



**Figure 4.** What do we want this figure to actually be?



**Figure 5.** Correlation between the optimal behaviour in the split-half and (a) adaptive choice paradigms; (b) average run length in the conjunction foraging paradigm; and (c) optimal behaviour in the adaptive choice paradigm and mean run length on the conjunction experiment.

with top-down visual search performance where task relevance is crucial, but not with bottom-up visual search tasks where salience in the environment guides attention [18]

Mention [19].

## References

- Walter R Boot, Arthur F Kramer, Ensar Becic, Douglas A Wiegmann, and Tate Kubose. Detecting transient changes in dynamic displays: The more you look, the less you see. *Human Factors*, 48(4):759–773, 2006.
- David H Brainard. The psychophysics toolbox. *Spatial vision*, 10:433–436, 1997.
- Alasdair DF Clarke, Patrick Green, Mike J Chantler, and Amelia R Hunt. Human search for a target on a textured background is consistent with a stochastic model. *Journal of vision*, 16(7):4–4, 2016.
- Frans W Cornelissen, Enno M Peters, and John Palmer. The eyelink toolbox: eye tracking with matlab and the psychophysics toolbox. *Behavior Research Methods, Instruments, & Computers*, 34(4):613–617, 2002.
- Marian Dawkins. Shifts of ‘attention’ in chicks during feeding.

distracting visual information [10,11,12]. Behavioural work in visual search has extended this research to demonstrate that working memory correlates only

- 464 *Animal Behaviour*, 19(3):575 – 582, 1971. 522
- 465 6. Iain D Gilchrist and Monika Harvey. 523
- 466 Evidence for a systematic component 525
- 467 within scan paths in visual search. 526
- 468 *Visual Cognition*, 14(4-8):704–715, 2006. 526
- 469 7. Jessica L Irons and Andrew B Leber. 527
- 470 Choosing attentional control settings in 528
- 471 dynamically changing environment. 529
- 472 *Attention, Perception, & Psychophysics*, 2016. 530
- 473 8. Jessica L Irons and Andrew B Leber. 531
- 474 Characterizing individual variation in the 532
- 475 strategic use of attentional control. 533
- 476 *Journal of Experimental Psychology: Human* 534
- 477 *Perception and Performance*, pages 1–18, 2018. 535
- 478 9. Ómar I. Jóhannesson, Árni Kristjánsson, 536
- 479 and Ian M. Thornton. 537
- 480 Are foraging patterns in humans related to 538
- 481 working memory and inhibitory control? 539
- 482 *Japanese Psychological Research*, 59(2):152– 540
- 483 166. 541
- 484 10. Ómar I Jóhannesson, Ian M Thornton, 542
- 485 Irene J Smith, Andrey Chetverikov, and 543
- 486 Árni Kristjánsson. 544
- 487 Visual foraging with fingers and eye gaze. 545
- 488 *i-Perception*, 7(2):2041669516637279, 2016. 546
- 489 11. Árni Kristjánsson, Ómar I Jóhannesson, 547
- 490 and Ian M Thornton. 548
- 491 Common attentional constraints in visual 549
- 492 foraging. 550
- 493 *PloS one*, 9(6):e100752, 2014. 551
- 494 12. Norman H Mackworth. 552
- 495 The breakdown of vigilance during 553
- 496 prolonged visual search. 554
- 497 *Quarterly Journal of Experimental Psychology*, 555
- 498 1(1):6–21, 1948. 556
- 499 13. Jiri Najemnik and Wilson S Geisler. 557
- 500 Eye movement statistics in humans are 558
- 501 consistent with an optimal search strategy. 559
- 502 *Journal of Vision*, 8(3):4–4, 2008. 560
- 503 14. Anna Nowakowska, Alasdair D F Clarke, 561
- 504 and Amelia R Hunt. 562
- 505 Human visual search behaviour is far from 563
- 506 ideal. 564
- 507 *Proceedings of the Royal Society B: Biological* 565
- 508 *Sciences*, 2017. 566
- 509 15. Anna Nowakowska, Alasdair DF Clarke, 567
- 510 Arash Sahraie, and Amelia R Hunt. 568
- 511 Inefficient search strategies in simulated 569
- 512 hemianopia. 570
- 513 *Journal of Experimental Psychology: Human* 571
- 514 *Perception and Performance*, 42(11):1858, 572
- 515 2016. 573
- 516 16. Anna Nowakowska, Alasdair D.F. Clarke, 574
- 517 Arash Sahraie, and Amelia R. Hunt. 575
- 518 Practice-related changes in eye movement 576
- 519 strategy in healthy adults with simulated 577
- 520 hemianopia. 578
- 521 *Neuropsychologia*, 2018. 579
- 522 17. Michael J Proulx. 580
- 523 Individual differences and metacognitive 581
- 524 knowledge of visual search strategy. 582
- 525 *PLoS One*, 6(10):e27043, 2011. 583
- 526 18. Kenith V Sobel, Matthew P Gerrie, Bradley J 584
- 527 Poole, and Michael J Kane. 585
- 528 Individual differences in working memory 586
- 529 capacity and visual search: The roles of top- 587
- 530 down and bottom-up processing. 588
- 531 *Psychonomic Bulletin & Review*, 14(5):840– 589
- 532 845, 2007. 590
- 533 19. Gijsbert Stoet. 591
- 534 Sex differences in search and gathering 592
- 535 skills. 593
- 536 *Evolution and Human Behavior*, 32(6):416– 594
- 537 422, 2011. 595
- 538 20. Anne M Treisman and Garry Gelade. 596
- 539 A feature-integration theory of attention. 597
- 540 *Cognitive psychology*, 12(1):97–136, 1980. 598
- 541 21. Edward K Vogel and Edward Awh. 599
- 542 How to exploit diversity for scientific gain: 600
- 543 Using individual differences to constrain 601
- 544 cognitive theory. 602
- 545 *Current Directions in Psychological Science*, 603
- 546 17(2):171–176, 2008. 604
- 547 22. Jeremy Wolfe, Matthew Cain, Krista 605
- 548 Ehinger, and Trafton Drew. 606
- 549 Guided search 5.0: Meeting the challenge of 607
- 550 hybrid search and multiple-target foraging. 608
- 551 *Journal of vision*, 15(12):1106–1106, 2015. 609